PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification ⁶:
C12N 15/12, C07K 14/475, A61K 37/02 A1

(11) International Publication Number:

WO 96/27007

(43) International Publication Date:

6 September 1996 (06.09.96)

(21) International Application Number:

PCT/AU96/00094

AU

ΑU

AU

(22) International Filing Date:

22 February 1996 (22.02.96)

(30) Priority Data:

Ć

PN 1457 2 March 1995 (02.03.95)
PN 6647 20 November 1995 (20.11.95)
PN 7274 22 December 1995 (22.12.95)

MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, ARIPO patent (KE, LS, MW, SD, SZ, UG), Eurasian patent (AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).

(81) Designated States: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IS,

JP, KE, KG, KP, KR, KZ, LK, LR, LS, LT, LU, LV, MD,

(71) Applicant (for all designated States except US): AMRAD OPERATIONS PTY. LTD. [AU/AU]; 17-27 Cotham Road, Kew, VIC 3101 (AU).

(72) Inventors; and

(75) Inventors/Applicants (for US only): HAYWARD, Nicholas, Kim [AU/AU]; 13 Prince Street, Paddington, QLD 4064 (AU). WEBER, Gunther [SE/AU]; 17-27 Cotham Road, Kew, VIC 3101 (AU). GRIMMOND, Sean [AU/AU]; 210 Swann Road, Taringa, QLD 4068 (AU). NORDENSKJOLD, Magnus [SE/SE]; Koltrastvagen 19, S-161 37 Stockholm (SE). LARSSON, Catharina [SE/SE]; Dannemoragatan 10m 3tr, S-113 44 Stockholm (SE).

(74) Agents: HUGHES, E., John, L. et al.; Davies Collison Cave, 1 Little Collins Street, Melbourne, VIC 3000 (AU). **Published**

With international search report.

(54) Title: A NOVEL GROWTH FACTOR AND A GENETIC SEQUENCE ENCODING SAME

(57) Abstract

6

The present invention relates generally to an isolated molecule having vascular endothelial growth factor-like properties and to a genetic sequence encoding same. The molecule will be useful in the development of a range of therapeutics and diagnostics useful in the treatment, prophylaxis and/or diagnosis of conditions requiring enhanced or diminished vasculature and/or vascular permeability. The molecule of the present invention is also a useful effector of primary and central neurons and is capable of inducing astroglial proliferation.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AM	Armenia	GB	United Kingdom	MW	Malawi
AT	Austria	GE	Georgia	MX	Manawi Mexico
AU	Australia	GN	Guinea	NE.	
BB	Barbados	GR	Greece		Niger
BE	Belgium	HU	Hungary	NL	Netherlands
BF	Burkina Faso	IE	Ireland	NO	Norway
BG	Bulgaria	IT.	Italy	NZ	New Zealand
BJ	Benin	JP	Japan	PL	Poland
BR	Brazil	KE	•	PT	Portugal
BY	Belarus	KG	Kenya	RO	Romania
CA	Canada	KP	Kyrgystan	RU	Russian Federation
CF	Central African Republic	A.F	Democratic People's Republic	SD	Sudan
CG	Congo	KR	of Korea	SE	Sweden
СН	Switzerland		Republic of Korea	SG	Singapore
CI	Côte d'Ivoire	KZ	Kazakhstan	SI	Slovenia
CM	Cameroon	u	Liechtenstein	SK	Slovakia
CN	China	LK	Sri Lanka	SN	Senegal
cs	Czechoslovakia	LR	Liberia	SZ	Swaziland
cz		LT	Lithuania	TD	Chad
DE	Czech Republic	LU	Luxembourg	TG	Togo
DK	Germany	LV	Larvia	TJ	Tajikistan
	Denmark	MC	Monaco	TT	Trinidad and Tobago
EE	Estonia	MD	Republic of Moldova	UA	Ukraine
ES	Spain	MG	Madagascar	UG	Uganda
FI	Finland	ML	Mali	US	United States of America
FR	France	MN	Mongolia	UZ	Uzbekistan
GA	Gabon	MR	Mauritania	VN	Viet Nam

30

A NOVEL GROWTH FACTOR AND A GENETIC SEQUENCE **ENCODING SAME**

The present invention relates generally to an isolated molecule having vascular endothelial growth factor-like properties and to a genetic sequence encoding same. The molecule will be useful in the development of a range of therapeutics and diagnostics useful in the treatment, prophylaxis and/or diagnosis of conditions requiring enhanced or diminished vasculature and/or vascular permeability. The molecule of the present invention is also a useful effector of primary and central neurons and is capable of 10 inducing astroglial proliferation.

Bibliographic details of the publications referred to by author in this specification are collected at the end of the description. Sequence Identity Numbers (SEQ ID NOs.) for 15 the nucleotide and amino acid sequences referred to in the specification are defined following the bibliography.

Throughout this specification, unless the context requires otherwise, the word "comprise", or variations such as "comprises" or "comprising", will be understood to imply the inclusion of a stated element or integer or group of elements or integers but not the exclusion of any other element or integer or group of elements or integers.

Vascular endothelial growth factor (hereinafter referred to as "VEGF"), also known as vasoactive permeability factor, is a secreted, covalently linked homodimeric glycoprotein that specifically activates endothelial tissues (Senger et al., 1993). A range of functions 25 have been attributed to VEGF such as its involvement in normal angiogensis including formation of the corpus luteum (Yan et al., 1993) and placental development (Sharkey et al., 1993), regulation of vascular permeability (Senger et al., 1993), inflammatory angiogenesis (Sunderkotter et al., 1994) and autotransplantation (Dissen et al., 1994) and human diseases such as turnour promoting angiogenesis (Folkman & Shing, 1992), rheumatoid arthritis (Koch et al., 1994) and diabetes related retinopathy (Folkman & Shing, 1992).

VEGF is, therefore, an important molecule making it a potentially valuable target for research into therapeutics, prophylactics and diagnostic agents based on VEGF or its activities. There is also a need to identify homologues or otherwise related molecules for use as an alternative to VEGF or in conjunction with VEGF.

5

10

In work leading up to the present invention, the inventors sought the multiple endocrine neoplasia type I susceptibility gene (MEN1). Surprisingly, the inventors discovered that a genetic sequence excluded as a candidate for the MEN1 gene was nevertheless a new growth factor having some similarity to VEGF. Furthermore, the growth factor of the present invention is an effector molecule for primary and central neurons.

Accordingly, one aspect of the present invention comprises a biologically isolated proteinaceous molecule comprising a sequence of amino acids which:

- (i) is at least about 15% similar to the amino acid sequence set forth in SEQ ID
 NO:2; and
 - (ii) is at least 5% dissimilar to the amino acid sequence set forth in SEQ ID NO:2.

Another aspect of the present invention provides a biologically isolated proteinaceous molecule having the following characteristics:

- 20 (i) comprises an amino acid sequence having at least about 15% similarity but at least about 5% dissimilarity to all or part of the amino acid sequence set forth in SEQ ID NO:2;
 - (ii) exhibits at least one property in common with VEGF.
- A related aspect of the present invention contemplates a biologically isolated proteinaceous molecule having the following characteristics:
 - (i) comprises an amino acid sequence having at least about 15% similarity but at least about 5% dissimilarity to the amino acid sequence set forth in SEQ ID NO:2:
- 30 (ii) exhibits at least one of the following properties:
 - (a) ability to induce proliferation of vascular endothelial cells;
 - (b) ability to interact with flt-1/flk-1 family of receptors;

20

25

ability to induce cell migration, cell survival and/or an increase in (c) intracellular levels of alkaline phosphatase.

By "biologically isolated" is meant that the molecule has undergone at least one step of 5 purification from a biological source. Preferably, the molecule is also biologically pure meaning that a composition comprises at least about 20%, more preferably at least about 40%, still more preferably at least about 65%, even still more preferably at least about 80-90% or greater of the molecule as determined by weight, activity or other convenient means, relative to other compounds in the composition. Most preferably, the molecule is sequencably pure.

Another preferred aspect of the present invention provides the molecule in recombinant form.

- According to this aspect of the present invention, there is provided a recombinant molecule comprising a sequence of amino acids which:
 - is at least about 15% similar to the amino acid sequence set forth in SEQ ID (i) NO:2; and
 - is at least 5% dissimilar to the amino acid sequence set forth in SEQ ID NO:2. (ii)

A related aspect of the present invention is directed to a recombinant molecule having the following characteristics:

- comprises an amino acid sequence having at least about 15% similarity but at (i) least about 5% dissimilarity to all or part of the amino acid sequence set forth in SEQ ID NO:2:
- (ii) exhibits at least one property in common with VEGF.

A further related aspect of the present invention contemplates a recombinant molecule having the following characteristics:

comprises an amino acid sequence having at least about 15% similarity but at 30 (i) least about 5% dissimilarity to the amino acid sequence set forth in SEQ ID NO:2;

WO 96/27007 PCT/AU96/00094

- 4 -

- (ii) exhibits at least one of the following properties:
 - (a) ability to induce proliferation of vascular endothelial cells;
 - (b) ability to interact with flt-1/flk-1 family of receptors;
 - (c) ability to induce cell migration, cell survival and/or an increase in intracellular levels of alkaline phosphatase.

The present invention also contemplates genomic or partial genome clones encoding a proteinaceous molecule having at least about 15% amino acid similarity but at least about 5% dissimilarity to SEQ ID NO:1.

10

15

20

5

The amino acid sequence set forth in SEQ ID NO:2 corresponds to human VEGF (referred to herein as "VEGF₁₆₅"). Accordingly, the molecule of the present invention is VEGF-like or is a homologue of VEGF but comprises an amino acid sequence which is similar but non-identical to the amino sequence of VEGF. Although the present invention is exemplified using a human VEGF-like molecule, this is done with the understanding that the instant invention contemplates the homologous molecule and encoding sequence from other mammals such as livestock animals (e.g. sheep, pigs, horses and cows), companion animals (e.g. dogs and cats) and laboratory test animals (e.g. mice, rats, rabbits and guinea pigs) as well as non-mammals such as birds (e.g. poultry birds), fish and reptiles. In a most preferred embodiment, the VEGF-like molecule is of human origin and encoded by a gene located at chromosome 11q13. The present invention extends, therefore, to the genomic sequence or part thereof encoding the subject VEGF-like molecule.

- Preferably, the percentage similarity is at least about 30%, more preferably at least about 40%, still more preferably at least about 50%, still even more preferably at least about 60-70%, yet even more preferably at least about 80-95% to all or part of the amino acid sequence set forth in SEQ ID NO:2.
- In a particularly preferred embodiment, the VEGF-like molecule of the present invention comprises a sequence of amino acids as set forth in SEQ ID NO:4 or is a part, fragment, derivative or analogue thereof. Particularly preferred similarities include about 19-20%,

30

and 29-30%. Reference herein to derivatives also includes splice variants. Accordingly, the present invention extends to splice variants of SOM175. Examples of splice variants contemplated by the present invention include but are not limited to variants with an amino acid sequence substantially as set forth in at least one of SEQ ID NO:6, SEQ ID NO:8 and/or SEQ ID NO:10 or mutants or derivatives or further splice variants thereof.

Another embodiment provides a recombinant molecule having the following characteristics:

- (i) an amino acid sequence substantially as set forth in SEQ ID NO:4 or having at least about 15% similarity to all or part thereof provided that said amino acid sequence is at least about 5% dissimilar to all or part of the amino acid sequence set forth in SEQ ID NO:2;
 - (ii) exhibits at least one biological property in common with VEGF.
- 15 Another embodiment provides a recombinant molecule having the following characteristics:
 - (i) an amino acid sequence substantially as set forth in SEQ ID NO:6 or having at least about 15% similarity to all or part thereof provided that said amino acid sequence is at least about 5% dissimilar to all or part of the amino acid sequence set forth in SEQ ID NO:2;
 - (ii) exhibits at least one biological property in common with VEGF.

Another embodiment provides a recombinant molecule having the following characteristics:

- an amino acid sequence substantially as set forth in SEQ ID NO:8 or having at least about 15% similarity to all or part thereof provided that said amino acid sequence is at least about 5% dissimilar to all or part of the amino acid sequence set forth in SEQ ID NO:2;
 - (ii) exhibits at least one biological property in common with VEGF.

Another embodiment provides a recombinant molecule having the following characteristics:

WO 96/27007 PCT/AU96/00094

- 6 -

(i) an amino acid sequence substantially as set forth in SEQ ID NO:10 or having at least about 15% similarity to all or part thereof provided that said amino acid sequence is at least about 5% dissimilar to all or part of the amino acid sequence set forth in SEQ ID NO:2;

5 (ii) exhibits at least one biological property in common with VEGF.

Such properties of VEGF include at least one of:

15

25

- (a) ability to induce proliferation of vascular endothelial cells;
- (b) an ability to interact with flt-1/flk-1 family of receptors;
- 10 (c) an ability to induce cell migration, cell survival and/or an increase in intracellular levels of alkaline phosphatase.

In accordance with the present invention, a preferred similarity is at least about 40%, more preferably at least about 50% and even more preferably at least about 65% similarity.

Still a further aspect of the present invention contemplates a peptide fragment corresponding to a portion of the amino acid sequence set forth in SEQ ID NO:4 or a splice variant thereof such as set forth in SEQ ID NO:6, SEQ ID NO:8 or SEQ ID NO:10 or a chemical equivalent thereof. The biologically isolated or recombinant molecule of the present invention may be naturally glycosylated or may comprise an altered glycosylation pattern depending on the cells from which it is isolated or synthesised. For example, if produced by recombinant means in prokaryotic organisms, the molecule would be non-glycosylated. The molecule may be a full length, naturally occurring form or may be a truncated or otherwise derivatised form.

Yet another aspect of the present invention is directed to a nucleic acid molecule encoding the VEGF-like molecule herein described. More particularly, the present invention provides a nucleic acid molecule comprising a sequence of nucleotides substantially as set forth in SEQ ID NO:3 or having at least 15% similarity to all or part thereof or being capable of hybridising under low stringency conditions to a reverse complement of the nucleotide sequence as set forth in SEQ ID NO:3 provided that the

15

20

nucleic acid sequence having at least 15% similarity but at least 30% dissimilarity to the nucleotide sequence as set forth in SEQ ID NO:3. The nucleotide sequence set forth in SEQ ID NO:3 is also referred to herein as "SOM175". Preferably, the percentage dissimilarity is about 35%, more preferably about 39% and even more preferably about 40-50% or greater.

For the purposes of defining the level of stringency, reference can conveniently be made to Sambrook *et al* (1989) at pages 9.47-9.51 which is herein incorporated by reference where the washing steps disclosed are considered high stringency. A low stringency is defined herein as being in 4-6X SSC/0.1-0.5% w/v SDS at 37-45°C for 2-3 hours. Depending on the source and concentration of nucleic acid involved in the hybridisation, alternative conditions of stringency may be employed such as medium stringent conditions which are considered herein to be 1-4X SSC/0.25-0.5% w/v SDS at \geq 45°C for 2-3 hours or high stringent conditions considered herein to be 0.1-1X SSC/0.1% w/v SDS at 60°C for 1-3 hours.

The present invention further contemplates a nucleic acid molecule which encodes a VEGF-like molecule as hereinbefore described having at least 15% nucleotide sequence homology to SEQ ID NO:3. Preferred levels of homology include at least about 40%, more preferably around 60-70%.

The present invention is further directed to the murine homologue of human VEGF (referred to herein as "mVRF"). The mVRF has approximately 85% identity and 92% conservation of amino acid residues over the entire coding region compared to human VEGF. The mVRF is encoded by a nucleic acid molecule comprising a nucleotide sequence substantially as set forth in Figure 9.

The VEGF-like molecule of the present invention will be useful in the development of a range of therapeutic and/or diagnostic applications alone or in combination with other molecules such as VEGF. The present invention extends, therefore, to pharmaceutical compositions comprising the VEGF-like molecule or parts, fragments, derivatives, homologues or analogues thereof together with one or more pharmaceutically acceptable

carriers and/or diluents. Furthermore, the present invention extends to vectors comprising the nucleic acid sequence set forth in SEQ ID NO:3 or having at least about 15%, more preferably about 40% and even more preferably around 60-70% similarity thereto but at least 30% and more preferably around 39% dissimilarity thereto and host cells comprising same. In addition, the present invention extends to ribozymes and antisense molecules based on SEQ ID NO:3 as well as neutralizing antibodies to the VEGF-like molecule. Such molecules may be useful in ameliorating the effects of, for example, over expression of VEGF-like genes leading to angiogenesis or vascularization of tumours.

10

15

30

Another aspect of the present invention contemplates a method of inducing astroglial proliferation in a mammal, said method comprising administering to said mammal an effective amount of a recombinant proteinaceous molecule having the characteristics:

- (i) comprises an amino acid sequence having at least about 15% similarity but at least about 5% dissimilarity to the sequence set forth in SEQ ID NO:2;
- (ii) exhibits at least one property in common with vascular endothelial growth factor (VEGF),

said administration being for a time and under conditions sufficient to induce astroglial proliferation.

Preferably, the recombinant proteinaceous molecule comprises the amino acid sequence set forth in SEQ ID NO:3 or SEQ ID NO:6.

- A further aspect of the present invention provides a method of promoting neural survival and/or proliferation in a mammal, said method comprising administering to said mammal an effective amount of a recombinant proteinaceous molecule having the characteristics:
 - (i) comprises an amino acid sequence having at least about 15% similarity but at least about 5% dissimilarity to the sequence set forth in SEQ ID NO:2;
 - (ii) exhibits at least one property in common with vascular endothelial growth factor (VEGF),

said administration being for a time and under conditions sufficient to induce astroglial proliferation.

Preferably, the recombinant proteinaceous molecule comprises the amino acid sequence set forth in SEQ ID NO:3 or SEQ ID NO:6.

The present invention also contemplates antibodies to the VEGF-like molecule or nucleic acid probes to a gene encoding the VEGF-like molecule which are useful as diagnostic agents.

10

The present invention is further described by reference to the following non-limiting Figures and/or Examples.

In the Figures:

15

- Figure 1 Nucleotide sequence [SEQ ID NO:1] and corresponding amino acid sequence [SEQ ID NO:2] of VEGF₁₆₅.
- Figure 2 Nucleotide sequence [SEQ ID NO:3] and corresponding amino acid 20 sequence [SEQ ID NO:4] of SOM175.
 - Figure 3 Results of BLAST search with SOM175 protein sequence.
 - Figure 4 BESTFIT alignment of VEGF cDNA and SOM175 cDNA.

25

- Figure 5 Multiple alignment of VEGF₁₆₅ with SOM175 and its splice variants at the nucleotide level.
- Figure 6 Multiple alignment of VEGF₁₆₅ with SOM175 and its splice variants at the amino acid level.
 - Figure 7 Diagrammatic representation of SOM175 and its splice variants.

WO 96/27007 PCT/AU96/00094

- 10 -

Figure 8(a) Diagrammatic representation of genomic structure of human SOM175 genomic showing exon/intron map.

Figure 8(b) Diagrammatic representation of genomic structure of human SOM175 showing exon/intron boundries.

Figure 9 Nucleotide and predicted peptide sequences derived from mVRF cDNA clones. Numbering of nucleotides are given on the left, starting from the A of the initiation codon. Amino acids are numbered on the right, starting from the first residue of the predicted mature protein after the putative signal peptide has been removed. The alternately spliced region is double underlined and the resulting peptide sequence from each mRNA is included. A potential polyadenylation signal is indicated in boldface. Start and stop codons of mVRF₁₆₇ and mVRF₁₈₆ are underlined and a polymorphic AC repeat in the 3' UTR is indicated by a stippled box. The positions of intron/exons boundaries are indicated by arrowheads.

Figure 10 BESTFIT alignments of human and murine VRF protein isoforms. A: mVRF₁₆₇ and hVRF₁₆₇. B: mVRF₁₈₆ and hVRF₁₈₆ from the point where the sequences diverge from the respective 167 amino acid isoforms. Amino acid identities are marked with vertical bars and conserved amino acids with colons. An arrow marks the predicted signal peptide cleavage site of human and mouse VRF.

15

20

Figure 11 BESTFIT alignment of mVRF₁₆₇ and mVEGF₁₈₈ (Breier *et al*, 1992) peptide sequences. An arrow marks the signal peptide cleavage site of mVEGF.

Identical amino acids are indicated by vertical bars and conservative substitutions by colons. Numbering of amino acids is as described in the legend to Figure 9.

Figure 12 Comparison of gene structure between VRF (a generic VRF gene is shown since the intron/exon organisation of the mouse and human homologues is almost identical) and other members of the human VEGF/PIGF/PDGF gene family. Exons are represented by boxes. Protein coding regions and untranslated regions are shown by filled and open sections respectively. The hatched region in VRF indicates the

additional 3' UTR sequence formed by alternate splicing of the VRF₁₈₆ isoform. Potential alternate splice products of each gene are shown.

- Figure 13 Autoradiogram of a Northern blot of total RNA from various adult mouse tissues (as indicated) hybridised with an mVRF cDNA clone. A major transcript of 1.3 kb was detected in all samples.
- Film autoradiographs (A-C) and dark-field micrographs (D-E) illustrating Figure 14 the expression pattern of mVRF and mRNA in the mouse. In the E14 mouse embryo (A) positive signals are present over the developing heart (Ha) and cerebral cortex (Cx). A low background signal is also present over other tissues in the section. In the E17 embryo (B) and the heart (Ha) is clearly visible due to a strong hybridisation signal. An equally strong signal is present over brown adipose tissue (Fa) in the back and around the thoracic cage. A moderate hybridisation signal is present over the spinal cord (SC) and the tongue (T). The background signal is reduced compared with the E14 embryo. In the young adult mouse (C-D), positive signals are present over the heart (Ha) and adipose tissue (Fa) around the thoracic cage, while, for example, the lungs (Lu) are unlabeled). The hybridisation signal over the heart is evenly distributed over the entire left ventricle, including papillary muscles (D). In the E17 heart hybridised with an excess of cold probe, no positive signal is present (E). Scale bars = 0.5 mm (A), 1.2 20 mm (B), 1 mm (C), 0.3 mm (D), 0.1 mm (E).
- Figure 15 Dark (A and C) and bright-field (B and D) micrographs showing mVRF mRNA expression in mouse adipose tissue (A-B) and spinal cord (C-D). A strong hybridisation signal is present over fat (A), as shown by the strong labeling in Sudan black stained sections (B). A weak signal is present also in skeletal muscle (M in A-B). In the adult spinal cord (C) the mVRF probes gave a neuronal staining pattern over the gray matter. Toloudine counterstaining showing that motoneurons in the ventral horn (D), interneurons in the deep part of the dorsal horn and around the central canal (not shown) where largely positive for mVRF mRNA. Scale bars = 0.1 mm (A), 0.1 mm (B), 0.25 mm (C), 0.015 mm (D).

Figure 16 Effect of VEGF on embryonic day 8 (E8) chick sensory neurons as determined by % survival, % neurite outgrowth and average neurite length (µm).

Figure 17 Effects of VEGF and SOM175 on chick glia. Tested were CNS glial, peripheral glia and CNS oligodendrocytes.

Figure 18 Effect of various SOM175 proteins on mouse astroglial cells.

A H (cpm)

- 1. FGF-2 (10 ng/ml) positive control
- 2. SOM_AX6^e 1 ng/ml
- 10 3. SOM_AX6 10 ng/ml
 - 4. SOM_AX6 100 ng/ml
 - 5. SOMAX6 1000 ng/ml
 - 6. SOMAX6 1000 ng/ml, no heparin
 - 7. SOMX6** 1 ng/ml
- 15 8. SOMX6 10 ng/ml
 - 9. SOMX6 100 ng/ml
 - 10. SOMX6 1000 ng/ml
 - 11. SOMX6 1000 ng/ml, no heparin
 - This refers to SOM175 absent exon 6;
- 20 ** This refers to SOM175.

25

Figure 19 Effect of various SOM175 proteins on mouse oligodenroglial cells. ■ ³H (cpm)

- 1. FGF-2 (10 ng/ml) positive control
- 2. SOM_AX6° 1 ng/ml
 - 3. SOMAX6 10 ng/ml
 - 4. SOMAX6 100 ng/ml
 - 5. SOMaX6 1000 ng/ml
 - 6. SOMAX6 1000 ng/ml, no heparin
- 30 7. SOMX6** 1 ng/ml
 - 8. SOMX6 10 ng/ml
 - 9. SOMX6 100 ng/ml,

- 10. SOMX6 1000 ng/ml
- 11. SOMX6 1000 ng/ml, no heparin
- This refers to SOM175 absent exon 6;
- This refers to SOM175.

Figure 20 Effect of various SOM175 proteins on mouse forebrain neurons. % survival

- 1. FGF-2 (10 ng/ml) positive control
- 2. SOM_AX6° 1 ng/ml
- 10 3. SOM_AX6 10 ng/ml
 - 4. SOMaX6 100 ng/ml
 - 5. SOM_aX6 1000 ng/ml
 - 6. SOMaX6 1000 ng/ml, no heparin
 - 7. SOMX6** 1 ng/ml
- 15 8. SOMX6 10 ng/ml
 - 9. SOMX6 100 ng/ml
 - 10. SOMX6 1000 ng/ml
 - 11. SOMX6 1000 ng/ml, no heparin
 - This refers to SOM175 absent exon 6;
- This refers to SOM175.

TABLE 1 - SUMMARY OF SEQUENCE IDENTITY NUMBERS

5		
	SEQ ID NO:1	Nucleotide sequence of VEGF ₁₆₅
	SEQ ID NO:2	Amino acid sequence of VEGF ₁₆₅
	SEQ ID NO:3	Nucleotide sequence of SOM175 (VEGF-like molecules)
	SEQ ID NO:4	Amino acid sequence of SOM175
10	SEQ ID NO:5	Nucleotide sequence of SOM175 absent exon 6
	SEQ ID NO:6	Amino acid sequence of SOM175 absent exon 6
	SEQ ID NO:7	Nucleotide sequence of SOM175 absent exon 6 and exon 7
	SEQ ID NO:8	Amino acid sequence of SOM175 absent exon 6 and exon 7
	SEQ ID NO:9	Nucleotide sequence of SOM175 absent exon 4
15	SEQ ID NO:10	Amino acid sequence of SOM175 absent exon 4
	SEQ ID NO:11	Oligonucleotide
	SEQ ID NO:12	Oligonucleotide
	SEQ ID NO:13	Oligonucleotide
	SEQ ID NO:14	Oligonucleotide
20		

EXAMPLE 1

Human cDNA clones

The original SOM175 cDNA was isolated by screening a human foetal brain library (λzapII, Stratagene) with the cosmid D11S750 (Larsson et al, 1992). The plasmid was excised "in vivo" and a single 1.1kb cDNA was obtained. Three independent SOM175 cDNAs clones were also isolated from a human foetal spleen library (Strategane, Unizap) using the above-mentioned SOM175 insert as a probe. Three clones were obtained: SOM175-4A, -5A and -6A. SOM175-5A is an alternately spliced clone with exon 4 being absent (SOM175-e4). These library screens were performed using hybridisation conditions recommended by the manufacturer of the library (Stratagene) and random primed insert of SOM175.

Two partial human SOM175 cDNAs have also isolated from a λGT11 human melanoma cell line A2058 library (Clontech) cDNA library screens were performed using hybridisation conditions described by Church and Gilbert, 1984). In each case, the probe was generated by random priming of a PCR product derived from SOM175 (18f-700r).

Mouse cDNA Clones

Human SOM175 was also used to screen a mouse neonatal whole brain cDNA library (Unizap, Stratagene). Four non-chimeric clones were isolated: M175-A, B, C, D. All clones were partial cDNAs and M175-C contained several introns. Three of these cDNAs lacked the exon 6.

Another clone referred to as M1 was completely sequenced and was found to contain the full open reading frame plus part of the 5'utr and total 3'utr.

15

EXAMPLE 2

DNA SEQUENCE ANALYSIS

The entire sequence of the cDNA clone (SOM175) was compiled and is shown in Figure 2 with its corresponding amino acid sequence. This sequence was screened for open reading frames using the MAP program (GCG, University of Wisconsin). A single open reading frame of 672bp was observed (see Figure 2). There appears to be little 5' untranslated sequences (2bp). The 3' untranslated region appears to be complete as it includes a poly-adenylation signal and poly-A tail.

Database homology searches were performed using the BLAST algorithm (run at NCBI, USA). This analysis revealed homology to several mammalian forms of VEGF (see Figure 3). The amount of homology between SOM175 and human VEGF₁₆₅ was determined using the BESTFIT program (GCG, University of Wisconsin; see Figures 4 and 5). Nucleotide homology was estimated at 69.7% and protein homology was estimated as at least 33.3% identity and 52.5% conservation using BESTFIT analysis. BLAST analysis on nucleotide sequences revealed the almost complete match to a human expressed sequence tag EST06302 (Adams et al., 1993).

These data indicate that SOM175 encodes a growth factor that has structural similarities to VEGF. Both genes show start and stop codons in similar positions and share discrete blocks of homology. All 8 cysteines as well as a number of other VEGF residues believed to be involved in dimerisation are conserved. These residues are Cysteine-47, Proline-70, Cysteine-72, Valine-74, Arginine-77, Cysteine-78, Glycine-80, Cysteine-81, Cysteine-82, Cysteine-89, Proline-91, Cysteine-122 and Cysteine-124 and are shown in Figure 6. Given the structural conservation between VEGF and the SOM175 gene product it is also possible that they share functional similarities. It is proposed that SOM175 encodes a VEGF-like molecule that shares some properties with VEGF but has unique properties of its own. The nucleotide sequence and corresponding amino acid sequence of VEGF₁₆₅ is shown in Figure 1.

EXAMPLE 3

The percentage similarity and divergence between VEGF₁₆₅ family and SOM175 family (protein) were analysed using the Clustal method, MegAlign Software, DNASTAR, Wisconsin. The results are shown in Tables 2.1 and 2.2. The alternatively spliced forms of SOM175 are abbreviated to SOM715-e6 where all of exon 6 is deleted; SOM715-e6 and 7 where all of exons 6 and 7 are deleted; and SOM175-e4 where all of exon 4 is deleted. The spliced form of SOM175 are shown in Figure 7. Genomic maps of SOM175 showing intron/exon boundaries are shown in Figure 8a and 8b.

Table 2.1

A Percent nucleotide similarity between splice variants of SOM175 and 5 human VEGF₁₆₅

		VEGF ₁₆₅	SOM175	SOM175-e6	SOM175-e6&7	SOM175-e4
	VEGF ₁₆₅	***	34.9	39.7	41.4	37.0
10	SOM175		***	98.9	95.1	99.2
	SOM175-e6			***	98.8	84.0
	SOM175-e6&7				***	80.3
	SOM175-e4					***

	VEGF ₁₆₅	SOM175	SOM175-e6	SOM175-e6&7	SOM175-e4
VEGF ₁₆₅	***	41.7	41.6	41.7	41.8
SOM175		***	0.2	0.2	0.0
SOM175-e6			***	0.0	0.2
SOM175-e6&7				***	0.3
SOM175-e4					***
	SOM175 SOM175-e6 SOM175-e6&7	VEGF ₁₆₅ *** SOM175 SOM175-e6 SOM175-e6&7	VEGF ₁₆₅ *** 41.7 SOM175 SOM175-e6 SOM175-e6&7	VEGF ₁₆₅ *** 41.7 41.6 SOM175 *** 0.2 SOM175-e6 SOM175-e6&7	VEGF ₁₆₅ **** 41.7 41.6 41.7 SOM175 **** 0.2 0.2 SOM175-e6 **** 0.0 SOM175-e6&7 ****

Table 2.2

15 A Percent amino acid identity between splice variants of SOM175 and human VEGF₁₆₅

		VEGF ₁₆₅	SOM175	SOM175-e6	SOM175-e6&7	SOM175-e4
20	VEGF ₁₆₅	***	31.4	42.3	33.5	40.6
	SOM175		***	74.7	73.7	99.1
	SOM175-e6			***	76.8	99.1
	SOM175-e6&7				***	99.1
	SOM175-e4					***
25	İ					

B Percent amino acid divergence between splice variants of SOM175 and human $VEGF_{165}$

5		VEGF ₁₆₅	SOM175	SOM175-e6	SOM175-e6&7	SOM175-e4
	VEGF ₁₆₅	***	65.7	55.4	54.6	57.4
	SOM175		***	19.9	4.2	0.0
	SOM175-e6			***	0.0	0.0
10	SOM175-e6&7				***	0.0
	SOM175-e4					***

15

EXAMPLE 4

BIOASSAYS TO DETERMINE THE FUNCTION OF SOM175

Assays are conducted to evaluate whether SOM175 has similar activities to VEGF on endothelial cell function, angiogenesis and wound healing. Other assays are performed based on the results of receptor binding distribution studies.

20

Assays of endothelial cell function

Endothelial cell proliferation. Endothelial cell growth assays as described in Ferrara & Henzel (1989) and in Gospodarowicz et al (1989).

25 Vascular permeability assay. This assay, which utilises the Miles test in guinea pigs, will be performed as described in Miles & Miles (1952).

Cell adhesion assay. The influence of SOM175 on adhesion of polymorphs to endothelial cells is analysed.

30

Chemotaxis. This is performed using the standard Boyden chamber chemotaxis assay.

Plasminogen activator assay. Endothelial cells are tested for plasminogen activator and plasminogen activator inhibitor production upon addition of SOM175 (Pepper et al (1991)).

5 Endothelial cell migration assay. The ability of SOM175 to stimulate endothelial cells to migrate and form tubes is assayed as described in Montesano et al (1986).

Angiogenesis Assav

SOM175 induction of an angiogenic response in chick chorioallantoic membrane is evaluated as described in Leung *et al* (1989).

Possible neurotrophic actions of SOM175 are assessed using the following assays:

Neurite outgrowth assay and gene induction (PC12 cells)

PC12 cells (a phaeochromocytoma cell line) respond to NGF and other neurotrophic factors by developing the characteristics of sympathetic neurons, including the induction of early and late genes and the extension of neurites. These cells are exposed to SOM175 and their response monitored (Drinkwater et al (1991); and Drinkwater et al (1993)).

20

Cultured neurons from the Peripheral Nervous System (PNS)

Primary cultures of the following PNS neurons are exposed to SOM175 and monitored for any response:

- sensory neurons from neural crest and dorsal root ganglia
- 25 sympathetic neurons from sympathetic chain ganglia
 - placode derived sensory neurons from nodose ganglia
 - motoneurons from spinal cord

The assays are described in Suter et al (1992) and in Marinou et al (1992).

Where an *in vitro* response is observed, *in vivo* assays for properties such as uptake and retrograde transport are performed as described in Hendry *et al* (1992).

Nerve regeneration (PNS)

Where neurotrophic effects of SOM175 are observed, its possible role in the regeneration of axotomised sensory neurons, sympathetic neurons and motoneurons is analysed by the methods of Otto et al (1989); Yip et al (1984) and Hendry et al (1976).

Actions of SOM175 on CNS neurons

The ability of SOM175 to promote survival of central nervous system neurons is analysed as described in Hagg et al (1992); Williams et al (1986); Hefti (1986) and Kromer (1987).

Wound Healing

10

The ability of SOM175 to support wound healing are tested in the most clinically relevant model available, as described in Schilling et al (1959) and utilised by Hunt et al (1967).

The Haemopoietic System

A variety of *in vitro* and *in vivo* assays on specific cell populations of the haemopoietic system are available and are outlined below:

20 Stem Cells

Murine

A variety of novel in vitro murine stem cell assays have been developed using FACS-purified cells:

25 (a) Repopulating Stem Cells

These are cells capable of repopulating the bone marrow of lethally irradiated mice, and have the Lin⁻, Rh^{hi}, Ly-6A/E⁺, c-kit⁺ phenotype. The test substance is tested on these cells either alone, or by co-incubation with multiple factors, followed by measurement of cellular proliferation by ³H thymidine incorporation.

(b) Late Stage Stem Cells

These are cells that have comparatively little bone marrow repopulating ability but can generate D13 CFU-S. These cells have the Lin-, Rhhi, Ly-6A/E+, c-kit+ phenotype. The test substance is incubated with these cells for a period of time, injected into lethally irradiated recipients, and the number of D13 spleen colonies enumerated.

(c) Progenitor-Enriched Cells

These are cells that respond in vitro to single growth factors, and have the Lin⁻, Rh^{hi}, Ly-6A/E⁺, c-kit⁺ phenotype. This assay will show if SOM175 can act directly on haemopoietic progenitor cells. The test substance is incubated with these cells in agar cultures, and the number of colonies enumerated after 7-14 days.

15 Atherosclerosis

Smooth muscle cells play a crucial role in the development or initiation of atherosclerosis, requiring a change in their phenotype from a contractile to a synthetic state. Macrophages, endothelial cells, T lymphocytes and platelets all play a role in the development of atherosclerotic plaques by influencing the growth and phenotypic modulations of smooth muscle cell. An *in vitro* assay that measures the proliferative rate and phenotypic modulations of smooth muscle cells in a multicellular environment is used to assess the effect of SOM175 on smooth muscle cells. The system uses a modified Rose chamber in which different cell types are seeded onto opposite coverslips.

25

30

20

Effects of SOM175 on bone

The ability of SOM175 to regulate proliferation of osteoblasts is assayed as described in Lowe et al (1991). Any effects on bone resorption are assayed as described in Lowe et al (1991). Effects on osteoblast migration and changes in intracellular molecules (e.g. cAMP accumulation, alkaline phosphatase levels) are analysed as described in Midy et al (1994).

Effects on skeletal muscle cells

Effects of SOM175 on proliferation of myoblasts and development of myotubes can be determined as described by Ewton et al (1980) and by Gospodarowicz et al (1976).

5

EXAMPLE 5 CLONING MURINE VEGF DNA

Isolation of cDNAs

Murine VRF (mVRF) clones were selected from a lambda Zap new born whole brain cDNA library (Stratagene). Primary phage from high density filters (5 \times 10⁴ pfu/plate) were identified by hybridisation with a 682bp ³²P-labelled probe generated by PCR from an hVRF cDNA (pSOM175) as described above. Hybridisation and stringent washes of nylon membranes (Hybond-N) were carried out at 65°C under conditions described by Church and Gilbert (1984). Positive plaques were picked, purified and excised in vivo to produce bacterial colonies containing cDNA clones in

Isolation of genomic clones

Genomic clones were isolated from a mouse strain SV/129 library cloned in the lambda Fix II vector (Stratagene). High density filters (5 x 10⁴ pfu/filter) were 20 screened with a 563 bp 32P-labelled probe generated by PCR amplification of the nucleotide 233-798 region of the mVRF cDNA (see Figure 9). Positive clones were plugged and re-screened with filters containing 400-800 pfu. Large scale phage preparations were prepared using the QIAGEN lambda kit or by ZnCl₂ purification 25 (Santos, 1991).

Nucleotide sequencing and analysis

cDNAs were sequenced on both strands using a variety of vector-based and internal primers with Applied Biosystems Incorporated (ABI) dye terminator sequencing kits according to the manufacturer's specifications. Sequences were analysed on an ABI Model 373A automated DNA sequencer. Peptide homology alignments were performed using the program BESTFIT (GCG, Wisconsin).

Identification of intron/exon boundaries

Identification of exon boundaries and flanking regions was carried out using PCR with mouse genomic DNA or mVRF genomic lambda clones as templates. The primers used in PCR to identify introns were derived from the hVRF sequence and to allow for potential human-mouse sequence mismatches annealing temperatures 5-10°C below the estimated T_m were used. All PCR products were sized by agarose gel electrophoresis and gel purified using QIAquick spin columns (Qiagen) and the intron/exon boundaries were sequenced directly from these products. In addition, some splice junctions were sequenced from subcloned genomic fragments of MVRF. Intron/exon boundaries were identified by comparing cDNA and genomic DNA sequences.

Northern analysis

Total cellular RNA was prepared from a panel of fresh normal adult mouse tisues

(brain, kidney, liver, muscle) using the method of Chomczynski and Sacchi (1987).

20µg of total RNA were electrophoresed, transferred to a nylon membrane (Hybond N, Amersham) and hybridised under standard conditions (Church & Gilbert, 1984).

Filters were washed at 65°C in 0.1xSSC (20xSSC is 3M NaCl/0.3M trisodium citrate), 0.1% SDS and exposed to X-ray film with intensifying screens at -70°C for

1-3 days.

Characterisation of mVRF cDNAs

Murine VRF homologues were isolated by screening a murine cDNA library with an hVRF cDNA clone. Five clones of sizes varying from 0.8-1.5 kb were recovered and sequenced. The cDNA sequences were complied to give a full length 1041 bp cDNA sequence covering the entire open reading frame (621 bp or 564 bp depending on the splice form, see below) and 3' UTR (379 bp), as well as 163 bp of the 5' UTR (Figure 9).

The predicted initiation codon matched the position of the start codon in hVRF. One other out of frame ATG was located at position -47 and two termination codons were observed upstream (positions -9 and -33, respectively) and in-frame with the putative

initiation codon.

The predicted N-terminal signal peptide of hVRF appears to be present in mVRF with 81% identity (17/21 amino acids). Peptide cleavage within mVRF is expected to occur after reside 21 (Figure 10). These data suggest that mature mVRF is secreted and could therefore conceivably function as a growth factor.

As with hVRF, two open reading frames (ORFs) were detected in cDNAs isolated by library screening. Four of five clones were found to be alternatively spliced and lacked a 101 bp fragment homologous to exon 6 of hVRF. The predicted peptide sequences of the two isoforms of mVRF were determined and aligned with the corresponding human isoforms (Figure 10).

The message encoding mVRF₁₈₆ contains a 621 bp ORF with coding sequences terminating at position +622, towards the end of exon 7 (Figure 9). The smaller message encoding mVRF₁₆₇ actually terminates downstream of the +622 TAG site due to a frame shift resulting from splicing out of the 101 bp exon 6 and the introduction of a stop codon (TGA) at position +666, near the beginning of exon 8 (Figure 9).

20

10

The mVRF₁₈₆ protein has strong homology to the amino and central portions of VEGF while the carboxyl end is completely divergent an is alanine rich. mVRF₁₆₇ possesses these similarities and also maintains homology to mVEGF right through to the C-terminus (Figure 11). The overall homology of mVRF₁₆₇ to hVRF₁₆₇ was 85% identity and 92% similarity, respectively (Figure 10). Likewise, homology between mVRF₁₆₇ and mVEGF (Breier *et al*, 1992) was 49% identity and 71% conservative amino acid substitution, respectively (Figure 11).

A canonical vertebrate polyadenylation signal (AATAAA) (Birnstiel et al, 1986) was not present in the mVRF cDNA, however, the closely matching sequence GATAAA is present at similar positions in both mouse and human VRF cDNAs (Figure 9). In contrast to hVRF, mVRF was found to contain an AC dinucleotide repeat at the

15

20

25

30

extreme 3' end of the 3' UTR (nucleotide positions 998 to 1011, Figure 9). Polymorphism of this repeat region was observed between some of the mVRF cDNAs, with the number of dinucleotides varying from 7 to 11.

5 Genomic characterisation of mVRF

Intron/exon boundaries (Table 3) were mapped using primers which flanked sequences homologous to the corresponding hVRF boundaries. Introns I, III, IV and VI of mVRF (Table 3, Figure 12) were smaller than the hVRF intervening sequences. The complete genomic sequence was compiled from the 5' UTR of mVRF through to intron VI, the largest intervening region (2.2 kb), by sequencing amplified introns and cloned genomic portions of mVRF. There was only one major difference in genomic structure between mVRF and hVRF and that was the exon 7/intron VI boundary of mVRF was located 10bp further downstream in relation to the cDNA sequence, hence exon 7 in mVRF is 10bp longer than the corresponding exon in hVRF.

Exons 6 and 7 are contiguous in mVRF, as has been found to occur in the human homologue. The strong sequence homology between exon 6 of mVRF and hVRF (Figure 10) suggests that this sequence is not a retained intronic sequence but rather encodes a functional part of the VRF₁₈₆ isoform.

General intron/exon structure is conserved between the various members of the VEGF gene family (VEGF, PIGF, hVRF) and therefore it is not surprising that the overall genomic organisation of the mVRF gene is very similar to these genes (Figure 12).

Previous comparative mapping studies have shown that the region surrounding the human multiple endocrine neoplasia type 1 disease locus on chromosome 11q13 is syntenic with the proximal segment of mouse chromosome 19 (Rochelle et al, 1992). Since the inventors have mapped the hVRF gene to within 1kb of the human MEN1 locus (see above) it is most likely that the murine VRF gene maps near the centromere of chromosome 19.

Expression studies of mVRF

Northern analysis of RNA from adult mouse tissues (muscle, heart, lung and liver) showed that expression appears to be ubiquitous and occurs primarily as a major band of approximately 1.3kb in size (Figure 14). This is somewhat different to the pattern observed for hVRF in which two major bands of 2.0 and 5.5 kb have been identified in all tissues examined. The 1.3 kb murine message presumably corresponds to the shorter of the human transcripts and the size variation thereof is most likely due to a difference in the length of the respective 5' UTRs.

10

15

EXAMPLE 6

EXPRESSION OF MURINE VEGF IN PRE- AND POST-NATAL MOUSE Animals

Timed pregnant (n=4) and young adult (n=2) mice (C57 inbred strain, ALAB, Sweden) were sacrificed with carbon dioxide, and the relevant tissues were taken out and frozen on a chuck. Tissues were kept at -70°C until further use. Two gestational ages was used in this study; embryonic day 8 (E8), 14 and £17.

In situ hybridisation histochemistry

In situ hybridisation was performed as previously described (Dagerlind et al, 1992).

- Briefly, transverse sections (14μm) were cut in a cryostat (Microm, Germany), thawed onto Probe-On slides (Fisher Scientific, USA) and stored in black sealed boxes at -70°C until used. The sequences of the synthetic 42-mer oligonucleotides complementary to mRNA encoding mVRF were
 - ACCACCACCTCCCTGGGCTGCATGTGGCACGTGCATAAACG [SEQ ID
- 25 NO:11] (complementary to nt 120-161) and
 AGTTGTTTGACCACATTGCCCATGAGTTCCATGCTCAGAGGC [SEQ ID
 NO:12] (complementary to nt 162-203). To detect the two alternative splice forms oligonucleotide GATCCTGGGGCTGGAGTGGATGGATGATGTCAGCTGG [SEQ ID NO:13] (complementary to nt xxx-xxx) and
- 30 GCGGGCAGAGGATCCTGGGGCTGTCTGGCCTCACAGCACT [SEQ ID NO:14] were used. The probes were labeled at the 3'-end with deoxyadenosine-alpha[thio]triphosphate [35S] (NEN, USA) using terminal deoxynucleotidyl

transferase (IBI, USA) to a specific activity of 7-10 x 108 cpm/µg and hybridised to the sections without pretreatment for 16-18 h at 42°C. The hybridisation mixture contained: 50% v/v formamide, $4 \times SSC$ (1 x SSC = 0.15M NaCl and 0.015M sodium-citrate), 1 x Denhardt's solution (0.02% each of polyvinyl-pyrrolidone, BSA and Ficoll), 1% v/v sarcosyl (N-lauroylsarcosine; Sigma), 0.02M phosphate buffer (pH 7.0), 10% w/v dextran sulfate (Pharmacia, Sweden), 250µg/ml yeast tRNA (Sigma), 500µg/ml sheared and heat denatured salmon sperm DNA (Sigma) and 200mM dithiothreitol (DTT; LKB, Sweden). In control sections, the specificity of both probes was checked by adding a 20-fold excess of unlabeled probe to the 10 hybridisation mixture. In addition, adjacent sections were hybridised with a probe unrelated to this study which gave a different expression pattern. Following hybridisation the sections were washed several times in 1 x SSC at 55°C, dehydrated in ethanol and dipped in NTB2 nuclear track emulsion (Kodak, USA). After 3-5 weeks the sections were development in D-19 developer (Kodak, USA) and coverslipped. In some cases, sections were opposed to an autoradiographic film (Beta-max 15 autoradiography film Amersham Ltd, UK) prior to emulsion-dipping.

The four different probes gave identical hybridisation patterns in all tissues examined. Mouse VRF expression was detecting already in the E8 embryo, in which positive signal was recorded over structures most likely corresponding to the neuronal tube. 20 In sagittal sections of E14 mouse embryo the strongest hybridisation signal was present over heart and in the nervous system, especially cerebral cortex (Figure 14A). A low level of expression was present in all other tissues. At a later gestational age, E17, a high mVRF mRNA signal was confined to he heart and brown fat tissue in the back and around the neck (Figure 14B). Clearly positive hybridisation signals 25 were present in the gray of the spinal cord and in the tongue (Figure 14B). Expression in the cerebral cortex was clearly reduced compared to day 14. The weak background expression seen in the E14 embryo in for example muscle, had decreased at this gestational age. A strong mVRF mRNA hybridisation signal was present solely over the heart and in the brown fat in the young adult mice (Figure 14C). The signal over the heart was evenly distributed ove the entire ventricular wall, including the papillary muscles (Figure 14D). In sections of heart tissue hybridised with an

15

excess of cold probe, no specific labeling over background signal was recorded (Figure 14E).

Apart from the heart, mVRF mRNA signal was present over certain tissues on the 5 outside of the thoracic cage that morphologically resembled brown fat. This was verified with sudan black counterstaining, which showed a strong staining in the same areas (Figure 15A and 15B). In transverse sections of adult mouse spinal cord, the mVRF probes gave a neuronal staining pattern over the gray matter (Figure 15C). Counterstaining with toluidine (Figure 15D) showed that motoneurons in the ventral horn (Figure 15C and 15D), interneurons (Figure 15C) in the deep part of the dorsal horn and around the central canal where to a large extent positive for mVRF mRNA.

EXAMPLE 7

EFFECTS OF VEGF AND SOM175 PROTEINS ON CHICK **SENSORY NEURONS**

The effects of VEGF and SOM175 proteins on embryonic day 8 chick sensory neurons were determined using the method of Nurcombe et al (1992). The neuronal assay was read at 48 hours using 2000 cells per assay well. The results were obtained using ³H-thymidine counts. The percentage survival of neurons, neurite 20 outgrowth and average neurite length in µm were determined using NGF as positive control and various concentrations of VEGF, VEGF in the presence of heparin and VEGF in the presence of heparin and 5 μ M, 5'-flurouracil (5FU). 5FU kills glial cells.

- The results are shown in Figure 16. The results show that VEGF is effective in 25 promoting neuronal survival but that this requires the presence of glial cells. Figure 17 shows the results of the effect of VEGF and SOM175 on three types of chick glia. The glia tested were CNS glia, peripheral glia and CNS oligodendrocytes. Heparin was used as 10 μg/ml in all cultures and the assay was read at 24 hours.
- Results were measured in ³H-thymidine counts using 2000 cells per well. 30

The results show that for chick central and peripheral neurons, astroglia were markedly stimulated to proliferate by SOM175 in the presence of heparin but that chick oligodendrocytes showed negligible increase in the rate of division.

5

30

EXAMPLE 8

EFFECTS OF SOM175 PROTEINS ON MOUSE PRIMARY AND CENTRAL NEURONS

The results in Example 7 show that VEGF isoform had an effect on chick primary and central neurons through the agency of the astroglial cells. Similar experiments were repeated in mouse cells.

Culture conditions

Neuronal and gligal cells for all *in vitro* experiments were prepared and cultured according o the techniques described in "Methods in Neurosciences (Vol. 2): Cell Culture" Ed. P.M. Conn, Academic Press, San Diego, 1990, pp33-46 for astroglial cells, pp56-74 for oligodendroglial cells, and pp87-102 for central neurons.

Cells were plated onto 24-well culture clusters (Nunc) coated with poly-L-ornithine

(0.1 mg/ml, 1h) at a density of 2,000 cells/well. After 48 hours in culture, neurons were counted in the wells under inverted phase light using well established techniques (Maruta et al. 1993) and glial cells assessed with [3H]thymidine uptake to monitor cell division rates as below. Heparin (10µg/ml, low molecular weight fraction, Sigma Chemical Corp.) was present at all times in the culture media except where noted. The neuronal cultures were supplemented with 5mM 5-fluoro-2-deoxyuridine (Sigma) to suppress background glial growth.

³H-Thymidine incorporation assay for glial cell proliferation

The cells were pulsed for 14h with ³H-thymidine (specific activity 103 µCi/ug) fraom a stock concentration of 0.1 mCi/ml in standard medium, giving a final incubating volume of 20 µl/well. The contents of the wells were harvested and absorbed onto nitrocellulose paper (Titertek, Flow). Remaining adherent cells were removed by

incubation with trypsin/versene (CSL Limited, Victoria, Australia) for 5 min. This procedure was carried out twice. The nitrocellulose discs were washed in a standard Titertek harvester (Flow) using first distilled water, and then methanol. The nitrocellulose discs were dried, scintillation fluid (containing 5% v/v Triton-X) added and the discs counted on a scintillation counter.

Greatest activity was seen with preparations of SOM175 absent exon 6 (SOMaX6) on mouse astroglial cell cultures, where there was a significant stimulus to their proliferation when delivered in conjunction with heparin (Figure 16). Little stimulus was given to the proliferation of oligodendroglial cells (Figure 17), and very little discernable potentiation of the survival response of isolated forebrain neurons (Figure 18). The standard deviation on all three graphs for each point was less than 8%.

The viability of neurons can be maintained by promoting glial cell proliferation.

5 Furthermore, SOMaX6 is a good inducer of astroglial proliferation and may be expressed in conjunction with the formation of astroglial endfeet on central nervous system endothelial cells.

Those skilled in the art will appreciate that the invention described herein is

susceptible to variations and modifications other than those specifically described. It
is to be understood that the invention includes all such variations and modifications.

The invention also includes all of the steps, features, compositions and compounds
referred to or indicated in this specification, individually or collectively, and any and
all combinations of any two or more of said steps or features.

TABLE 3
Splice junctions of the murine VRF gene

5	5' UTR*	Exon 1 >223bp	CCCAGgtacgtgcgt	Intron I	495bp
	ttccccacagGCCCC	Exon 2 43bp	GAAAGgtaataatag	Intron II	288bp
	ctgcccacagTGGTG	Exon 3 197bp	TGCAGgtaccagggc	Intron III	196bp
	ctgagcacagATCCT	Exon 4 74bp	TGCAGgtgccagccc	Intron IV	182bp
	ctcttttcagACCTA	Exon 5 36bp	GACAGattcttggtg	Intron V	191bp
10	ctcctcctagGGTTG	Exon 6 101bp		(no intron)	
	CCCACTCCAGCCC	CA Exon 7 135bp	TGTAGgtaaggagtc	Intron VI ~	-2200bp
	cactccccagGTGCC	Exon 8 394bp	AGAGATGGAGAC	ACT	

Uppercase and lowercase letters denote exonic and intronic sequences respectively.

15 * Indicates that the 5' end of exon 1 has not yet been determined.

BIBLIOGRAPHY

Adams MD, Soares MB, Kerlavage AR, Fields C, Venter JC, (1993) Nature Genet, 4, 373-380.

Birnstiel ML, Busslinger M and Strub K (1985) Cell 41, 349-359.

Breier G, Albrecht U, Sterrer S and Risau W (1992) Development 114, 521-532.

Chomczynski P and Sacchi N (1987) Analyt. Biochem. 162, 156-159.

Church G and Gilbert W (1984) Proc. Natl. Acad. Sci. USA 18, 1991-1995.

Dagerlind A, Friberg K, Bean AJ and Hokfelt T (1992) Histochemistry 98, 39-49.

Dissen GA, Lara HE, Fabrenbach WH, Costa ME, Ojeda SR, (1994) Endocrinology 134, 1146-1154.

Drinkwater CC, Barker PA, Suter U and Shooter EM (1993) J. Biol. Chem., 268, 23202-23207.

Drinkwater CC, Suter U, Angst C and Shooter EM (1991) Proc. Roy. Soc. Lond. (Series B). 246, 307-313.

Ewton DZ & Florini JR (1980) Endocrinology, 106: 577-583.

Ferrara N & Henzel WJ (1989) Biochem. Biophys. Res. Commun. 161, 851-858.

Folkman J & Shing Y (1992) J. Biol. Chem. 267, 10931-10934.

Gospodarowicz D, Abraham JA & Schilling J (1989) Proc. Natl. Acad. Sci USA 86, 7311-7315.

Gospodarowicz D, Weseman J, Morgan JS & Lindstrom J (1976) J. Cell Biol., 70: 395-405.

Hagg T, Quon D, Higaki J & Varon S (1992) Neuron, 8, 145-158.

Hefti S (1986) J. Neurosci, 6, 2155-2162.

Hendry IA & Campbell J (1976) J. Neurocytol., 5, 351-360.

Hendry IA, Murphy M, Hilton DJ, Nicola NA & Bartlett PF (1992) J. Neurosci. 12, 3427-3434.

Hunt et al., (1967) Am. J. Surgery, 114: 302-307.

Koch AE, Harlow LA, Haines GK, Amento EP, Unemoti EN, Wong WL, Pope RM, Ferrara N, (1994) J. Immunol. 152, 4149-4156.

Kromer AF (1987) Science, 235, 214-216.

Larsson C, Weber G, Kvanta E, Lewis C, Janson M, Jones C, Glaser T, Evans G, Nordenskjold M, (1992) Hum. Genet. 89, 187-193.

OZUS Leung DW, Cachianes G, Kuang W-J, Goeddel DV & Ferrara N (1989) Science 246:1306-1309.

Lowe C, Cornish J, Callon K, Martin TJ & Reid IR (1991) J. Bone Mineral Res., 6, 1277-1283.

Lowe C, Cornish J, Martin TJ & Reid IR (1991) Calcif. Tissue Int., 49, 394-397.

Martinou JC, Martinou I & Kato AC (1992) Neuron, 8, 737-744.

Maruta et al (1993) Growth Factors 8: 119-134.

Midy V & Plouet J (1994) Biochem. Biophys. Res. Commun., 199: 380-386.

Miles AA & Miles EM (1952) J. Physiol. (Lond) 118:228-257.

Montesano R, Vassalli JD, Baird A, Guillemin R & Orci, L (1986) Proc. Natl. Acad. Sci USA, 83, 7297-7301.

Nurcombe et al (1992) Development 116: 1175-1183.

Otto D., Frotscher M & Unsicker K (1989) J. Neurosci. Res., 22, 83-91.

Pepper MS, Ferrara N, Orci L, Montesano R. (1991) Biochem. Biophys. Res. Commun. 181, 902-906).

Rochell JM, Watson ML, Oakey RJ and Seldin MF (1992) Genomics 14, 26-31.

Roth S & Weston J (1967) Proc. Natl. Acad. Sci USA, 58: 974-980.

Sambrook J, Fritsch EF, Maniatis T, (1989) Molecular Cloning: A Laboratory Manual - 2nd Ed. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y.

Santos MA (1991) Nucleic Acids Res. 19, 5442.

Schilling et al., (1959) Surgery, 46: 702-710.

Senger DR, Van De Water L, Brown LF, Nagy JA, Yeo KT, Yeo TK, Berse B,

Jackman RW, Dvorak AM, Dvorak HF (1993) Cancer Netastasis Rev. 12, 303-324.

Sharkey AM, Chamock-Jones DS, Boocock CA, Brown KD, Smith SK, (1993) J. Reprod. Fertil. 99, 609-615.

Sunderkotter C, Steinbrink K, Goebeler M, Bhardway R, Sorg E, (1993) J. Leukocyt, Biol. 55, 410-422.

Suter U, Angst C, Tien C-L, Drinkwater CC, Lindsay RM and Shooter EM (1992) J. Neurosci., 12, 306-318.

Tischer E, Mitchell R, Hartman T, Silva M, Gospodarowicz D, Fiddes JC, & Abraham J (1991) J. Biol. Chem. 266, 11947-11954.

Williams LR, Varon S, Peterson GM, Wictorin K, Fischer W, Bjorklund A & Gage FH (1986) Proc. Natl. Acad. Sci. USA 83, 9231-9235.

Yan Z, Weich HA, Bernart W, Breckwoldt M, Neulen J, (1993) J. Clin. Endocrinol. Metab. 77, 1723-1725.

Yip NK, Rich KM, Lampe PA & Johnson EM Jr (1984) J. Neurosci., 4, 2986-2992.

SEQUENCE LISTING

(1) GENERAL INFORMATION:

(i) APPLICANT:

(countries other than US) AMRAD OPERATIONS PTY. LTD. (us only) Hayward, N and Weber, G

- (ii) TITLE OF INVENTION: A NOVEL GROWTH FACTOR AND A GENETIC SEQUENCE ENCODING SAME
- (iii) NUMBER OF SEQUENCES: 14
- (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: DAVIES COLLISON CAVE
 - (B) STREET: 1 LITTLE COLLINS STREET
 - (C) CITY: MELBOURNE
 - (D) STATE: VICTORIA
 - (E) COUNTRY: AUSTRALIA
 - (F) ZIP: 3000
- (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: PCT INTERNATIONAL
 - (B) FILING DATE: 22-FEB-1996
- (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: AU PN1457
 - (B) FILING DATE: 02-MAR-1995
 - (A) APPLICATION NUMBER: AU PN6647
 - (B) FILING DATE: 20-NOV-1995
 - (A) APPLICATION NUMBER: AU PN7274
 - (B) FILING DATE: 22-DEC-1995

(viii) ATTORNEY/AGENT INFORMATION: (A) NAME: HUGHES DR, E JOHN L (C) REFERENCE/DOCKET NUMBER: EJH/EK	
(ix) TELECOMMUNICATION INFORMATION: (A) TELEPHONE: +61 3 9254 2777 (B) TELEFAX: +61 3 9254 2770	
(2) INFORMATION FOR SEQ ID NO:1:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 649 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(ii) MOLECULE TYPE: DNA	
(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 17589	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:	
TCGGGCCTCC GAAACC ATG AAC TTT CTG CTG TCT TGG GTG CAT TGG AGC Met Asn Phe Leu Leu Ser Trp Val His Trp Ser 1 5 10	49
CTT GCC TTG CTG CTC TAC CTC CAC CAT GCC AAG TGG TCC CAG GCT GCA Leu Ala Leu Leu Tyr Leu His His Ala Lys Trp Ser Gln Ala Ala 15 20 25	97
CCC ATG GCA GAA GGA GGG GGG CAG AAT CAT CAC GAA GTG GTG AAG TTC Pro Met Ala Glu Gly Gly Gln Asn His His Glu Val Val Lys Phe 30 35 40	145
ATG GAT GTC TAT CAG CGC AGC TAC TGC CAT CCA ATC GAG ACC CTG GTG Met Asp Val Tyr Gln Arg Ser Tyr Cys His Pro Ile Glu Thr Leu Val 50 55	193
GAC ATC TTC CAG GAG TAC CCT GAT GAG ATC GAG TAC ATC TTC AAG CCA Asp Ile Phe Gln Glu Tyr Pro Asp Glu Ile Glu Tyr Ile Phe Lys Pro 60 65 70 75	241
TCC TGT GTG CCC CTG ATG CGA TGC GGG GGC TGC TGC AAT GAC GAG GGC Ser Cys Val Pro Leu Met Arg Cys Gly Cys Cys Arg	289

Ser Cys Val Pro Leu Met Arg Cys Gly Gly Cys Cys Asn Asp Glu Gly

50

CTG GAG TGT GTG CCC ACT GAG GAG TCC AAC ATC ACC ATG CAG ATT ATG Leu Glu Cys Val Pro Thr Glu Glu Ser Asn Ile Thr Met Gln Ile Met 95 100 105	337
CGG ATC AAA CCT CAC CAA GGC CAG CAC ATA GGA GAG ATG AGC TTC CTA Arg Ile Lys Pro His Gln Gly Gln His Ile Gly Glu Met Ser Phe Leu 110 115 120	385
CAG CAC AAC AAA TGT GAA TGC AGA CCA AAG AAA GAT AGA GCA AGA CAA Gln His Asn Lys Cys Glu Cys Arg Pro Lys Lys Asp Arg Ala Arg Gln 125 130 135	433
GAA AAT CCC TGT GGG CCT TGC TCA GAG CGG AGA AAG CAT TTG TTT GTA Glu Asn Pro Cys Gly Pro Cys Ser Glu Arg Arg Lys His Leu Phe Val 140 145 150 155	481
CAA GAT CCG CAG ACG TGT AAA TGT TCC TGC AAA AAC ACA GAC TCG CGT Gln Asp Pro Gln Thr Cys Lys Cys Ser Cys Lys Asn Thr Asp Ser Arg 160 165 170	529
TGC AAG GCG AGG CAG CTT GAG TTA AAC GAA CGT ACT TGC AGA TGT GAC Cys Lys Ala Arg Gln Leu Glu Leu Asn Glu Arg Thr Cys Arg Cys Asp 175 180 185	577
AAG CCG AGG CGG TGAGCCGGGC AGGAGGAAGG AGCCTCCCTC AGCGTTTCGG Lys Pro Arg Arg 190	629
GAACCAGATC TCTCACCAGG	649
(2) INFORMATION FOR SEQ ID NO:2:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 191 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: protein	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:	
Met Asn Phe Leu Leu Ser Trp Val His Trp Ser Leu Ala Leu Leu Leu 1 5 10 15	
Tyr Leu His His Ala Lys Trp Ser Gln Ala Ala Pro Met Ala Glu Gly 20 25 30	
Gly Gly Gln Asn His His Glu Val Val Lys Phe Met Asp Val Tyr Gln	

Arg Ser Tyr Cys His Pro Ile Glu Thr Leu Val Asp Ile Phe Gln Glu

60

Tyr 65	Pro	Asp	Glu	Ile	Glu 70	Туг	Ile	: Phe	. Lys	Pro		Cys	Val	Pro	Leu 80	
Met	Arg	Cys	Gly	Gly 85	Сув	Сув	Asn	Asp	Glu 90		' Leu	Glu	Cys	Val 95	Pro	
Thr	Glu	Glu	Ser 100	Asn	Ile	Thr	Met	Gln 105	Ile	Met	Arg	Ile	Lys 110	Pro	His	
Gln	Gly	Gln 115	His	Ile	Gly	Glu	Met 120	Ser	Phe	Leu	Gln	His 125	Asn	Lys	Сув	
Glu	Cys 130	Arg	Pro	Lys	Lys	As p 135	Arg	Ala	Arg	Gln	Glu 140	Asn	Pro	Cys	Gly	
Pro 145	Cys	Ser	Glu	Arg	Arg 150	Lys	His	Leu	Phe	Val 155	Gln	Asp	Pro	Gln	Thr 160	
Cys 1	Lys	Cys	Ser	Cys 165	Lys	Asn	Thr	Asp	Ser 170	Arg	Cys	Lys	Ala	Arg 175	Gln	
Leu (3lu	Leu	Asn 180	Glu .	Arg '	Thr	Cys	Arg 185	Cys .	Asp	Lys		Arg 190	Arg		
()	(i) iii) iix)	SEQU (A) (B) (C) (D) MOLE FEAT (A) (B)	UENCE TYPE TOF CCULE URE: NAM LOC	E CHAINGTH: PE: MANDE POLOG TYP E/KE ATIO	ARACT 105 Aucle DNES EY: 1 PE: D Y: C N: 3	TERIS 4 ba ic a is: a inea NA DS 62	STIC: ase pacid sing) ar	pairs le	NO:							
1	Se	r Pro	o Lei	ı Let	ı Arg	J Ar	J Le	u Lei	u Let	ı Ala	a Ala	a Le	u Le	u Gli	n 5	4
CTG GC Leu Al	a Pı	o Al	La GI	G GC .n Al	c cc a Pr	T GT	C TC	er Gl	G CC ln Pr 25	T GA	T GC	C CC	ro G	C CAG Ly Hi 30	C is	95
CAG AGG	G AA	's va	G GT 1 Va 5	G TC	A TG(AT) P Il	e As	r GTG Sp Va	TAT	ACI	CGC	g Al	ACC	TGC or Cy	'S	143

CAG CCC CGG GAG GTG GTG CCC TTG ACT GTG GAG CTC ATG GGC ACC Gln Pro Arg Glu Val Val Pro Leu Thr Val Glu Leu Met Gly Thr 50 55 60	191
GTG GCC AAA CAG CTG GTG CCC AGC TGC GTG ACT GTG CAG CGC TGT GGT Val Ala Lys Gln Leu Val Pro Ser Cys Val Thr Val Gln Arg Cys Gly 65 70 75	239
GGC TGC TGC CCT GAC GAT GGC CTG GAG TGT GTG CCC ACT GGG CAG CAC Gly Cys Cys Pro Asp Asp Gly Leu Glu Cys Val Pro Thr Gly Gln His 80 85 90 95	287
CAA GTC CGG ATG CAG ATC CTC ATG ATC CGG TAC CCG AGC AGT CAG CTG Gln Val Arg Met Gln Ile Leu Met Ile Arg Tyr Pro Ser Ser Gln Leu 100 105 110	335
GGG GAG ATG TCC CTG GAA GAA CAC AGC CAG TGT GAA TGC AGA CCT AAA Gly Glu Met Ser Leu Glu Glu His Ser Gln Cys Glu Cys Arg Pro Lys 115 120 125	383
AAA AAG GAC AGT GCT GTG AAG CCA GAC AGG GCT GCC ACT CCC CAC CAC Lys Lys Asp Ser Ala Val Lys Pro Asp Arg Ala Ala Thr Pro His His 130 135 140	431
CGT CCC CAG CCC CGT TCT GTT CCG GGC TGG GAC TCT GCC CCC GGA GCA Arg Pro Gln Pro Arg Ser Val Pro Gly Trp Asp Ser Ala Pro Gly Ala 145 150 155	479
CCC TCC CCA GCT GAC ATC ACC CAT CCC ACT CCA GCC CCA GGC CCC TCT Pro Ser Pro Ala Asp Ile Thr His Pro Thr Pro Ala Pro Gly Pro Ser 160 165 170 175	527
GCC CAC GCT GCA CCC AGC ACC AGC GCC CTG ACC CCC GGA CCT GCC Ala His Ala Ala Pro Ser Thr Thr Ser Ala Leu Thr Pro Gly Pro Ala 180 185 190	5 75
GCT GCC GCT GCC GCA GCT TCC TCC GTT GCC AAG GGC GGG GCT T Ala Ala Ala Asp Ala Ala Asa Ser Ser Val Ala Lys Gly Gly Ala 195 200 205	624
AGAGCTCAAC CCAGACACCT GCAGGTGCCG GAAGCTGCGA AGGTGACACA TGGCTTTTCA	684
GACTCAGCAG GGTGACTTGC CTCAGAGGCT ATATCCCAGT GGGGGAACAA AGGGGAGCCT	744
GGTAAAAAAC AGCCAAGCCC CCAAGACCTC AGCCCAGGCA GAAGCTGCTC TAGGACCTGG	804
GCCTCTCAGA GGGCTCTTCT GCCATCCCTT GTCTCCCTGA GGCCATCATC AAACAGGACA	864
GAGTTGGAAG AGGAGACTGG GAGGCAGCAA GAGGGGTCAC ATACCAGCTC AGGGGAGAAT	924
GGAGTACTGT CTCAGTTTCT AACCACTCTG TGCAAGTAAG CATCTTACAA CTGGCTCTTC	984
CTCCCCTCAC TAAGAAGACC CAAACCTCTG CATAATGGGA TTTGGGCTTT GGTACAAGAA	1044
CTGTGACCCC CAACCCTGAT AAAAGAGATG GAAGGAAAAA AAAAAAAAA	1004

- (2) INFORMATION FOR SEQ ID NO:4:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 207 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:
- Met Ser Pro Leu Leu Arg Arg Leu Leu Leu Ala Ala Leu Leu Gln Leu 1 5 10 15
- Ala Pro Ala Gln Ala Pro Val Ser Gln Pro Asp Ala Pro Gly His Gln
 20 25 30
- Arg Lys Val Val Ser Trp Ile Asp Val Tyr Thr Arg Ala Thr Cys Gln
 35 40 45
- Pro Arg Glu Val Val Pro Leu Thr Val Glu Leu Met Gly Thr Val
 50 55 60
- Ala Lys Gln Leu Val Pro Ser Cys Val Thr Val Gln Arg Cys Gly Gly 65 70 75 80
- Cys Cys Pro Asp Asp Gly Leu Glu Cys Val Pro Thr Gly Gln His Gln
 85 90 95
- Val Arg Met Gln Ile Leu Met Ile Arg Tyr Pro Ser Ser Gln Leu Gly
 100 105 110
- Glu Met Ser Leu Glu Glu His Ser Gln Cys Glu Cys Arg Pro Lys Lys 115 120 125
- Lys Asp Ser Ala Val Lys Pro Asp Arg Ala Ala Thr Pro His His Arg
- Pro Gln Pro Arg Ser Val Pro Gly Trp Asp Ser Ala Pro Gly Ala Pro 145 150 155 160
- Ser Pro Ala Asp Ile Thr His Pro Thr Pro Ala Pro Gly Pro Ser Ala 165 170 175
- His Ala Ala Pro Ser Thr Thr Ser Ala Leu Thr Pro Gly Pro Ala Ala 180 185 190
- Ala Ala Asp Ala Ala Ala Ser Ser Val Ala Lys Gly Gly Ala 195 200 205

(2)	INFORMATION	rok	SEQ	ID	NO:5:

(i)	SEQUENCE	CHARACTERISTICS:
	_	

- (A) LENGTH: 993 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA
- (ix) FEATURE:

130

- (A) NAME/KEY: CDS
- (B) LOCATION: 3..566
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

cc	ATG Met 1	AGC Ser	CCT Pro	CTG Leu	CTC Leu 5	CGC Arg	CGC Arg	CTG Leu	CTG Leu	CTC Leu 10	GCC Ala	GCA (Ala	CTC (Leu	TG C	AG Gln 15	47
CTC	GCC Ala	CCC Pro	GCC Ala	CAG Glr 20	ı Ala	CCI Pro	GTC Val	TCC Sea	CAG CGlr 25	Pro	GAT Asp	GCC Ala	CCT Pro	GGC Gly 30	CAC His	95
CAG Gln	AGG Arg	AAA Lys	GTG Val	Val	TCA Ser	TGG Trp	ATA Ile	GAT Asp	Val	TAT Tyr	ACT Thr	CGC (Arg	CT A Ala 45	CC T	GC Cys	143
CAG Gln	CCC Pro	CGG Arg 50	Glu	GTG Val	GTG Val	GTG Val	CCC Pro 55	Leu	ACT Thr	GTG (GAG (CTC A Leu 60	TG G Met	GC A	CC Thr	191
GTG Val	GCC Ala 65	AAA Lys	CAG Gln	CTG Leu	GTG Val	CCC Pro 70	AGC Ser	TGC Cys	GTG . Val	ACT (GTG (Val 75	CAG C	GC T Arg	GT G Cys	GT Gly	239
GGC Gly 80	TGC Cys	TGC Cys	CCT Pro	GAC Asp	GAT Asp 85	GGC Gly	CTG Leu	GAG Glu	TGT (Cys	GTG (Val 90	CCC # Pro	CT G Thr	GG C Gly	AG CA Gln	AC His 95	287
CAA Gln	GTC Val	CGG Arg	ATG Met	CAG Gln 100	ATC Ile	CTC . Leu	ATG : Met	ATC (CGG : Arg 105	TAC (CCG A	GC A	Ser	AG CT Gln : 110	G Leu	335
GGG Gly	GAG Glu	ATG Met	TCC Ser 115	CTG (Leu	GAA Glu	GAA (Glu	CAC / His	AGC (Ser 120	CAG 1	Cys	AA T Glu	GC AC	GA CO Arg	T AA Pro 1	A Lys	383

AAA AAG GAC AGT GCT GTG AAG CCA GAT AGC CCC AGG CCC CTC TGC CCA

135

Lys Lys Asp Ser Ala Val Lys Pro Asp Ser Pro Arg Pro Leu Cys Pro

CGC TGC ACC CAG CAC CAG CGC CCT GAC CCC CGG ACC TGC CGC TGC Arg Cys Thr Gln His His Gln Arg Pro Asp Pro Arg Thr Cys Arg Cys 145 150 155	479
CGC TGC CGA CGC CGC AGC TTC CTC CGT TGC CAA GGG CGG GGC TTA GAG Arg Cys Arg Arg Arg Ser Phe Leu Arg Cys Gln Gly Arg Gly Leu Glu 160 165 170 175	527
CTC AAC CCA GAC ACC TGC AGG TGC CGG AAG CTG CGA AGG TGACACATGG Leu Asn Pro Asp Thr Cys Arg Cys Arg Lys Leu Arg Arg 180 185	576
CTTTTCAGAC TCAGCAGGGT GACTTGCCTC AGAGGCTATA TCCCAGTGGG GGAACAAAGG	636
GGAGCCTGGT AAAAAACAGC CAAGCCCCCA AGACCTCAGC CCAGGCAGAA GCTGCTCTAG	696
GACCTGGGCC TCTCAGAGGG CTCTTCTGCC ATCCCTTGTC TCCCTGAGGC CATCATCAAA	756
CAGGACAGAG TTGGAAGAGG AGACTGGGAG GCAGCAAGAG GGGTCACATA CCAGCTCAGG	816
GGAGAATGGA GTACTGTCTC AGTTTCTAAC CACTCTGTGC AAGTAAGCAT CTTACAACTG	876
GCTCTTCCTC CCCTCACTAA GAAGACCCAA ACCTCTGCAT AATGGGATTT GGGCTTTGGT	936
ACAAGAACTG TGACCCCCAA CCCTGATAAA AGAGATGGAA GGAAAAAAAA AAAAAAA	993

(2) INFORMATION FOR SEQ ID NO:6:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 188 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

Met Ser Pro Leu Leu Arg Arg Leu Leu Leu Ala Ala Leu Leu Gln Leu 1 5 10 15

Ala Pro Ala Gln Ala Pro Val Ser Gln Pro Asp Ala Pro Gly His Gln
20 25 30

Arg Lys Val Val Ser Trp Ile Asp Val Tyr Thr Arg Ala Thr Cys Gln 35 40 45

Pro Arg Glu Val Val Val Pro Leu Thr Val Glu Leu Met Gly Thr Val 50 55 60

Ala Lys Gln Leu Val Pro Ser Cys Val Thr Val Gln Arg Cys Gly Gly 65 70 75 80

Cys Cys Pro Asp Asp Gly Leu Glu Cys Val Pro Thr Gly Gln His G 85 90 95	ln
Val Arg Met Gln Ile Leu Met Ile Arg Tyr Pro Ser Ser Gln Leu Gl 100 105 110	Y
Glu Met Ser Leu Glu Glu His Ser Gln Cys Glu Cys Arg Pro Lys Ly 115 120 125	's
Lys Asp Ser Ala Val Lys Pro Asp Ser Pro Arg Pro Leu Cys Pro Ar 130 135 140	g
Cys Thr Gln His His Gln Arg Pro Asp Pro Arg Thr Cys Arg Cys Ar 145 150 155 16	_
Cys Arg Arg Ser Phe Leu Arg Cys Gln Gly Arg Gly Leu Glu Le 165 170 175	u
Asn Pro Asp Thr Cys Arg Cys Arg Lys Leu Arg Arg 180 185	
(2) INFORMATION FOR SEQ ID NO:7: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 858 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA (ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 3431 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:	
CC ATG AGC CCT CTG CTC CGC CGC CTG CTG CTC GCC GC	47
CTG GCC CCC GCC CAG GCC CCT GTC TCC CAG CCT GAT GCC CCT GGC CAC Leu Ala Pro Ala Gln Ala Pro Val Ser Gln Pro Asp Ala Pro Gly His 20 25 30	95
CAG AGG AAA GTG GTG TCA TGG ATA GAT GTG TAT ACT CGC GCT ACC TGC Gln Arg Lys Val Val Ser Trp Ile Asp Val Tyr Thr Arg Ala Thr Cys 35 40 45	143
CAG CCC CGG GAG GTG GTG GTG CCC TTG ACT GTG GAG CTC ATG GGC ACC	191

Gln Pro Arg Glu Val Val Pro Leu Thr Val Glu Leu Met Gly Thr

55

Val Ala Lys Gln Leu Val Pro Ser Cys Val Thr Val Gln Ary Cys Gly 65 70 75	239
GGC TGC TGC CCT GAC GAT GGC CTG GAG TGT GTG CCC ACT GGG CAG CAC Gly Cys Cys Pro Asp Asp Gly Leu Glu Cys Val Pro Thr Gly Gln His 80 85 90 95	287
CAA GTC CGG ATG CAG ATC CTC ATG ATC CGG TAC CCG AGC AGT CAG CTG Gln Val Arg Met Gln Ile Leu Met Ile Arg Tyr Pro Ser Ser Gln Leu 100 105 110	335
GGG GAG ATG TCC CTG GAA GAA CAC AGC CAG TGT GAA TGC AGA CCT AAA Gly Glu Met Ser Leu Glu Glu His Ser Gln Cys Glu Cys Arg Pro Lys 115 120 125	383
AAA AAG GAC AGT GCT GTG AAG CCA GAT AGG TGC CGG AAG CTG CGA AGG Lys Lys Asp Ser Ala Val Lys Pro Asp Arg Cys Arg Lys Leu Arg Arg 130 135 140	431
TGACACATGG CTTTTCAGAC TCAGCAGGGT GACTTGCCTC AGAGGCTATA TCCCAGTGGG	491
GGAACAAAGG GGAGCCTGGT AAAAAACAGC CAAGCCCCCA AGACCTCAGC CCAGGCAGAA	551
GCTGCTCTAG GACCTGGGCC TCTCAGAGGG CTCTTCTGCC ATCCCTTGTC TCCCTGAGGC	611
CATCATCAAA CAGGACAGAG TTGGAAGAGG AGACTGGGAG GCAGCAAGAG GGGTCACATA	671
CCAGCTCAGG GGAGAATGGA GTACTGTCTC AGTTTCTAAC CACTCTGTGC AAGTAAGCAT	731
CTTACAACTG GCTCTTCCTC CCCTCACTAA GAAGACCCAA ACCTCTGCAT AATGGGATTT	791
GGGCTTTGGT ACAAGAACTG TGACCCCCAA CCCTGATAAA AGAGATGGAA GGAAAAAAA	851
AAAAAA	950

- (2) INFORMATION FOR SEQ ID NO:8:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 143 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:
- Met Ser Pro Leu Leu Arg Arg Leu Leu Leu Ala Ala Leu Leu Gln Leu 1 5 10 15
- Ala Pro Ala Gln Ala Pro Val Ser Gln Pro Asp Ala Pro Gly His Gln
 20 25 30
- Arg Lys Val Val Ser Trp Ile Asp Val Tyr Thr Arg Ala Thr Cys Gln
 35 40 45
- Pro Arg Glu Val Val Pro Leu Thr Val Glu Leu Met Gly Thr Val 50 55 60
- Ala Lys Gln Leu Val Pro Ser Cys Val Thr Val Gln Arg Cys Gly Gly 65 70 75 80
- Cys Cys Pro Asp Asp Gly Leu Glu Cys Val Pro Thr Gly Gln His Gln
 85 90 95
- Val Arg Met Gln Ile Leu Met Ile Arg Tyr Pro Ser Ser Gln Leu Gly
 100 105 110
- Glu Met Ser Leu Glu Glu His Ser Gln Cys Glu Cys Arg Pro Lys Lys 115 120 125
- Lys Asp Ser Ala Val Lys Pro Asp Arg Cys Arg Lys Leu Arg Arg 130 135 140

(2) INFORMATION	FOR	SEQ	ID	NO:	9	:
-----------------	-----	-----	----	-----	---	---

(i)	SEQUENCE	CHARACTERISTICS.

- (A) LENGTH: 910 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA

(ix) FEATURE:

- (A) NAME/KEY: CDS
- (B) LOCATION: 3..305

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

CC ATG AGC CCT CTG CTC CGC CGC CTG CTG CTC GCC GC	47
CTG GCC CCC GCC CAG GCC CCT GTC TCC CAG CCT GAT GCC CCT GGC CAC Leu Ala Pro Ala Gln Ala Pro Val Ser Gln Pro Asp Ala Pro Gly His 20 25 30	95
CAG AGG AAA GTG GTG TCA TGG ATA GAT GTG TAT ACT CGC GCT ACC TGC Gln Arg Lys Val Val Ser Trp Ile Asp Val Tyr Thr Arg Ala Thr Cys 35 40 45	143
CAG CCC CGG GAG GTG GTG CCC TTG ACT GTG GAG CTC ATG GGC ACC Gln Pro Arg Glu Val Val Val Pro Leu Thr Val Glu Leu Met Gly Thr 50 55 60	191
GTG GCC AAA CAG CTG GTG CCC AGC TGC GTG ACT GTG CAG CGC TGT GGT Val Ala Lys Gln Leu Val Pro Ser Cys Val Thr Val Gln Arg Cys Gly 65 70 75	239
GGC TGC TGC CCT GAC GAT GGC CTG GAG TGT GTG CCC ACT GGG CAG CAC Gly Cys Cys Pro Asp Asp Gly Leu Glu Cys Val Pro Thr Gly Gln His 80 85 90 95	287
CAA GTC CGG ATG CAG ACC TAAAAAAAAA GACAGTGCTG TGAAGCCAGA Gln Val Arg Met Gln Thr 100	335
CAGGGCTGCC ACTCCCCACC ACCGTCCCCA GCCCCGTTCT GTTCCGGGCT GGGACTCTGC	395
CCCCGGAGCA CCCTCCCCAG CTGACATCAC CCATCCCACT CCAGCCCCAG GCCCCTCTGC	455
CCACGCTGCA CCCAGCACCA CCAGCGCCCT GACCCCCGGA CCTGCCGCTG CCGCTGCCGA	515
CGCCGCAGCT TCCTCCGTTG CCAAGGGCGG GGCTTAGAGC TCAACCCAGA CACCTGCAGG	575
TGCCGGAAGC TGCGAAGGTG ACACATGGCT TTTCAGACTC AGCAGGGTGA CTTGCCTCAG	635
AGGCTATATC CCAGTGGGGA ACAAAGAGGA GCCTGGTAAA AAACAGCCAA GCCCCCAAGA	695

Val Arg Met Gln Thr

100

755

815

875

910

- 48 -
CCTCAGCCCA GGCAGAAGCT GCTCTAGGAC CTGGGCCTCT CAGAGGGCTC TTCTGCCATC
CCTTGTCTCC CTGAGGCCAT CATCAAACAG GACAGAGTTG GAAGAGGAGA CTGGGAGGCA
GCAAGAGGGG TCACATACCA GCTCAGGGGA GAATGGAGTA CTGTCTCAGT TTCTAACCAC
TCTGTGCAAG TAAGCATCTT ACAACTGGCT CTTCC
(2) INFORMATION FOR SEQ ID NO:10:
(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 101 amino acids(B) TYPE: amino acid(D) TOPOLOGY: linear
(ii) MOLECULE TYPE: protein
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:
Met Ser Pro Leu Leu Arg Arg Leu Leu Leu Ala Ala Leu Leu Gln Leu 1 5 10 15
Ala Pro Ala Gln Ala Pro Val Ser Gln Pro Asp Ala Pro Gly His Gln 20 25 30
Arg Lys Val Val Ser Trp Ile Asp Val Tyr Thr Arg Ala Thr Cys Gln

40

55

Pro Arg Glu Val Val Pro Leu Thr Val Glu Leu Met Gly Thr Val

Ala Lys Gln Leu Val Pro Ser Cys Val Thr Val Gln Arg Cys Gly Gly

Cys Cys Pro Asp Asp Gly Leu Glu Cys Val Pro Thr Gly Gln His Gln

(2) INFORMATION FOR SEQ ID NO::	:11:	NO:	ID	SEO	FOR	INFORMATION	(2)
---------------------------------	------	-----	----	-----	-----	-------------	-----

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 42 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Oligonucleotide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

ACCACCACCT CCCTGGGCTG GCATGTGGCA CGTGCATAAA CG

42

- (2) INFORMATION FOR SEQ ID NO:12:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 42 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Oligonucleotide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

AGTTGTTTGA CCACATTGCC CATGAGTTCC ATGCTCAGAG GC

42

- (2) INFORMATION FOR SEQ ID NO:13:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 38 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Oligonucleotide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:

GATCCTGGGG CTGGAGTGGG ATGGATGATG TCAGCTGG

- (2) INFORMATION FOR SEQ ID NO:14:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 40 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Oligonucleotide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

GCGGGCAGAG GATCCTGGGG CTGTCTGGCC TCACAGCACT

CLAIMS:

- 1. A biologically isolated proteinaceous molecule having the following characteristics:
 - (i) comprises an amino acid sequence having at least about 15% similarity but at least about 5% dissimilarity to the sequence set forth in SEQ ID NO:2;
 - (ii) exhibits at least one property in common with vascular endothelial growth factor (VEGF).
- 2. A proteinaceous molecule according to claim 1 wherein the molecule exhibits at least one of the following properties:
 - (i) an ability to induce vascular endothelial cells;
 - (ii) an ability to interact with flt-1/flk-1 family of receptors; and/or
 - (iii) an ability to induce cell migration, cell survival and/or an increase in intracellular lavels of alkaline phosphatase.
- 3. A proteinaceous molecule according to claim 1 or 2 wherein said molecule has the capacity to induce astroglial proliferation.
- 4. A proteinaceous molecule according to claim 1 wherein said molecule is of human origin.
- 5. A proteinaceous molecule according to claim 1 wherein said molecule is of non-human origin.
- 6. A proteinaceous molecule according to claim 5 wherein said molecule is of livestock animal, companion animal, laboratory test animal, avian, fish or reptilian origin.
- 7. A proteinaceous molecule according to claim 5 wherein said molecule is encoded by a gene located at chromosome 11q13.

- 8. A proteinaceous molecule according to claim 1 wherein the percentage similarity to SEQ ID NO:2 is at least about 30%.
- 9. A proteinaceous molecule according to claim 1 wherein the percentage similarity to SEQ ID NO:2 is at least about 40%.
- 10. A proteinaceous molecule according to claim 1 wherein the percentage similarity to SEQ ID NO:2 is at least about 60-70%.
- 11. A proteinaceous molecule according to claim 1 comprising a sequence of amino acids as set forth in SEQ ID NO:4 or a part, fragment, derivative or analogue thereof.
- 12. A proteinaceous molecule according to claim 1 comprising an amino acid sequence substantially set forth in SEQ ID NO:6 or a part, fragment, derivative or analogue thereof.
- 13. A proteinaceous molecule according to claim 1 comprising an amino acid sequence substantially set forth in SEQ ID NO:8 or a part, fragment, derivative or analogue thereof.
- 14. A proteinaceous molecule according to claim 1 comprising an amino acid sequence substantially set forth in SEQ ID NO:10 or a part, fragment, derivative or analogue thereof.
- 15. A recombinant molecule having the following characteristics:
 - an amino acid sequence substantially as set forth in SEQ ID NO:4
 or having at least about 15% similarity to but at least about 5%
 dissimilarity to the amino acid sequence set forth in SEQ ID
 NO:2;
 - (ii) exhibits at least one biological property in common with VEGF.

- 16. A recombinant molecule having the following characteristics:
 - (i) an amino acid sequence substantially as set forth in SEQ ID NO:6
 or having at least about 15% similarity to but at least about 5%
 dissimilarity to the amino acid sequence set forth in SEQ ID
 NO:2;
 - (ii) exhibits at least one biological property in common with VEGF.
- 17. A recombinant molecule having the following characteristics:
 - (i) an amino acid sequence substantially as set forth in SEQ ID NO:8 or having at least about 15% similarity to but at least about 5% dissimilarity to the amino acid sequence set forth in SEQ ID NO:2;
 - (ii) exhibits at least one biological property in common with VEGF.
- 18. A recombinant molecule having the following characteristics:
 - (i) an amino acid sequence substantially as set forth in SEQ ID NO:10 or having at least about 15% similarity to but at least about 5% dissimilarity to the amino acid sequence set forth in SEQ ID NO:2;
 - (ii) exhibits at least one biological property in common with VEGF.
- 19. A recombinant molecule according to claim 15 or 16 or 17 or 18 having at least one of the following properties:
 - (a) an ability to induce vascular endothelial cells;
 - (b) an ability to interact with flt1/flki family of receptors;
 - (c) an ability to induce cell migration, cell survival and/or increase intracellular levels of alkaline phosphatase.
- 20. A recombinant molecule according to claim 15 or 16 or 17 or 18 having the capacity to induce astroglial proliferation.

- 21. A recombinant molecule according to claim 20 wherein the molecule comprises an amino acid sequence substantially as set forth in SEQ ID NO:6.
- 22. A peptide fragment corresponding to a portion of the amino acid sequence set forth in SEQ ID NO:4 or a derivative or chemical equivalent thereof.
- 23. A peptide fragment according to claim 22 having the sequence set forth in SEQ ID NO:6 or a chemical equivalent thereof.
- 24. A peptide fragment according to claim 22 having the sequence set forth in SEQ ID NO:8 or a chemical equivalent thereof.
- 25. A peptide fragment according to claim 22 having the sequence set forth in SEQ ID NO:10 or a chemical equivalent thereof.
- 26. A nucleic acid molecule comprising a sequence of nucleotides or complementary to a sequence encoding a proteinaceous molecule having the following characteristics:
 - (i) comprises an amino acid sequence having at least about 15% similarity but at least about 5% dissimilarity to the sequence set forth in SEQ ID NO:2;
 - (ii) exhibits at least one property in common with vascular endothelial growth factor (VEGF).
- 27. A nucleic acid molecule according to claim 26 wherein the proteinaceous molecule exhibits at least one of the following properties:
 - (i) an ability to induce vascular endothelial cells;
 - (ii) an ability to interact with flt-1/flk-1 family of receptors; and/or
 - (iii) an ability to induce cell migration, cell survival and/or an increase in intracellular lavels of alkaline phosphatase.
- 28. A nucleic acid molecule according to claim 27 wherein the proteinaceous molecule has the capacity to induce astroglial proliferation.

- 29. A nucleic acid molecule according to claim 28 wherein said molecule encodes an amino acid sequence substantially as set forth in SEQ ID NO:6.
- 30. A nucleic acid molecule according to claim 1 wherein said molecule is of human origin.
- 31. A nucleic acid molecule according to claim 1 wherein the percentage similarity to SEQ ID NO:2 is at least about 30%.
- 32. A nucleic acid molecule according to claim 26 comprising a nucleotide sequence substantially as set forth in SEQ ID NO:3 or having at least 15% similarity thereto or capable of hybridising under low stringency conditions to a reverse complement of the nucleotide sequence as set forth in SEQ ID NO:3 provided that the nucleotide sequence has at least 15% similarity but at least 30% dissimilarity to the nucleotide sequence set forth in SEQ ID NO:3.
- 33. A nucleic acid molecule according to claim 26 encoding a murine homologue of human VEGF and comprising a nucleotide sequence substantially as set forth in Figure 9.
- 34. A pharmaceutical composition comprising a proteinaceous molecule according to claim 1 or 2 or 3 or 11 and one or more pharmaceutically acceptable carriers and/or diluents.
- 35. A method for preparing a recombinant molecule having the following characteristics:
 - (i) comprises an amino acid sequence having at least about 15% similarity but at least about 5% dissimilarity to the sequence set forth in SEQ ID NO:2;
 - (ii) exhibits at least one property in common with vascular endothelial growth factor (VEGF),

said method comprising expressing a nucleic acid molecule encoding said recombinant

molecule by a suitable host grown under conditions effective to synthesise said recombinant molecule and then isolating said molecule.

- 36. A method according to claim 35 wherein the nucleic acid molecule comprises a sequence of nucleotides as set forth in SEQ ID NO:3 or having at least 15% similarity thereto or is capable of hybridising under low stringency conditions to a reverse complement of the nucleotide sequence as set forth in SEQ ID NO:3 provided that the nucleotide sequence has at least 15% similarity but at least 30% dissimilarity to the nucleotide sequence set forth in SEQ ID NO:3.
- 37. A method of inducing astroglial proliferation in a mammal, said method comprising administering to said mammal an effective amount of a recombinant proteinaceous molecule having the characteristics:
 - comprises an amino acid sequence having at least about 15% similarity but at least about 5% dissimilarity to the sequence set forth in SEQ ID NO:2;
 - (ii) exhibits at least one property in common with vascular endothelial growth factor (VEGF),

said administration being for a time and under conditions sufficient to induce astroglial proliferation.

- 38. A method according to claim 37 wherein the recombinant proteinaceous molecule comprises an amino acid sequence substantially as set forth in SEQ ID NO:3 or is a derivative thereof.
- 39. A method according to claim 37 wherein the recombinant proteinaceous molecule comprises an amino acid sequence substantially as set forth in SEQ ID NO:6 or is a derivative thereof.
- 40. A method of promoting neuronal survival and/or proliferation in a mammal, said method comprising administering to said mammal an effective amount of a recombinant proteinaceous molecule having the characteristics:

- (i) comprises an amino acid sequence having at least about 15% similarity but at least about 5% dissimilarity to the sequence set forth in SEQ ID NO:2;
- (ii) exhibits at least one property in common with vascular endothelial growth factor (VEGF),

said administration being for a time and under conditions sufficient to induce astroglial proliferation.

- 41. A method according to claim 40 wherein the recombinant proteinaceous molecule comprises an amino acid sequence substantially as set forth in SEQ ID NO:3 or is a derivative thereof.
- 41. A method according to claim 40 wherein the recombinant proteinaceous molecule comprises an amino acid sequence substantially as set forth in SEQ ID NO:6 or is a derivative thereof.

2/52	3/52
Fig.1(i)	Fig.1(ii)
4/52	5/52
Fig.1(iii)	Fig.1(iv)

1	TCGGCCTCC GAAACC ATG AAC TTT CTG Met Asn Phe Leu 1
50	CTT GCC TTG CTG CTC TAC CTC CAC Leu Ala Leu Leu Tyr Leu His 15
98	CCC ATG GCA GAA GGA GGA GGG CAG Pro Met Ala Glu Gly Gly Gln 30
146	ATG GAT GTC TAT CAG CGC AGC TAC Met Asp Val Tyr Gln Arg Ser Tyr 45
194	GAC ATC TTC CAG GAG TAC CCT GAT Asp Ile Phe Gln Glu Tyr Pro Asp 60 65
242	TCC TGT GTG CCC CTG ATG CGA TGC Ser Cys Val Pro Leu Met Arg Cys 80
290	CTC GAG TGT GTG CCC ACT GAG GAG Leu Glu Cys Val Pro Thr Glu Glu 95
338	CGG ATC AAA CCT CAC CAA GGC CAG Arg Ily Lys Pro His Gln Gly Gln 110

Fig.1(i)

CTG TCT TGG GTG CAT TGG AGC Leu Ser Trp Val His Trp Ser 5	49
CAT GCC AAG TGG TCC CAG GCT GCA His Ala Lys Trp Ser Gln Ala Ala 20 25	97
AAT CAT CAC GAA GTG GTG AAG TTC Asn His His Glu Val Val Lys Phe 40	145
TGC CAT CCA ATC GAG ACC CTG GTG Cys His Pro Ile Glu Thr Leu Val	193
GAG ATC GAG TAC ATC TTC AAG CCA Glu Ile Glu Tyr Ile Phe Lys Pro 70 75	241
GGG GGC TGC TGC AAT GAC GAG GGC Gly Gly Cys Cys Asn Asp Glu Gly 85	289
TCC AAC ATC ACC ATG CAG ATT ATG Ser Asn Ile Thr Met Gln Ile Met 100	337
CAC ATA GGA GAG ATG AGC TTC CTA His Ile Gly Glu Met Ser Phe Leu 120	385

Fig.1(ii)

386	CAG CAC AAC AAA TGT GAA TGC AGA Gln His Asn Lys Cys Glu Cys Arg 125
434	GAA AAT CCC TGT GGG CCT TGC TCA Glu Asn Pro Cys Gly Pro Cys Ser 140
482	CAA GAT CCG CAG ACG TGT AAA TGT Gln Asp Pro Gln Thr Cys Lys Cys 160
530	TGC AAG GCG AGG CAG CTT GAG TTA Cys Lys Ala Arg Gln Leu Glu Leu 175
578	AAG CCG AGG CGG TGAGCCGGGC AGGAG Lys Pro Arg Arg 190
630	GAACCAGATC TCTCACCAGG

Fig.1(iii)

CCA AAG AAA GAT AGA GCA AGA CAA Pro Lys Lys Asp Arg Ala Arg Gln 135	433
GAG CGG AGA AAG CAT TTG TTT GTA Glu Arg Arg Lys His Leu Phe Val 150	481
TCC TGC AAA AAC ACA GAC TCG CGT Ser Cys Lys Asn Thr Asp Ser Arg 165	529
AAC GAA CGT ACT TGC AGA TGT GAC Asn Glu Arg Thr Cys Arg Cys Asp 180	577
GAAGG AGCCTCCCTC AGCGTTTCGG	629
	649

Fig.1(iv)

7/52	8/52
Fig.2(i)	Fig. 2 (ii)
9/52	10/52
Fig 2(iii)	Fig 2(iv)
11/52	. 12/52
Fig 2(v)	Fig 2(vi)

1	CC ATG AGC CCT CTG CTC CGC CGC Met Ser Pro Leu Leu Arg Arg
48	CTG GCC CCC GCC CAG GCC CCT GTC Leu Ala Pro Ala Gln Ala Pro Val 20
96	CAG AGG AAA GTG GTG TCA TGG ATA Gln Arg Lys Val Val Ser Trp Ile 35
144	CAG CCC CGG GAG GTG GTG CCC Gln Pro Arg Glu Val Val Val Pro 50
192	GTG GCC AAA CAG CTG GTG CCC AGC Val Ala Lys Gln Leu Val Pro Ser 65 70
240	GGC TGC TGC CCT GAC GAT GGC CTG Gly Cys Cys Pro Asp Asp Gly Leu 80
288	CAA GTC CGG ATG CAG ATC CTC ATG Gln Val Arg Met Gln Ile Leu Met 100
336	GGG GAG ATG TCC CTG GAA GAA CAC Gly Glu Met Ser Leu Glu Glu His 115

Fig.2(i)

CTG CTG CTC GCC GCA CTC CTG CAG Leu Leu Leu Ala Ala Leu Leu Gln 10	47
TCC CAG CCT GAT GCC CCT GGC CAC Ser Gln Pro Asp Ala Pro Gly His 25	95
GAT GTG TAT ACT CGC GCT ACC TGC Asp Val Tyr Thr Arg Ala Thr Cys 40	143
TTG ACT GTG GAG CTC ATG GGC ACC Leu Thr Val Glu Leu Met Gly Thr	191
TGC GTG ACT GTG CAG CGC TGT GGT Cys Val Thr Val Gln Arg Cys Gly 75	239
GAG TGT GTG CCC ACT GGG CAG CAC Glu Cys Val Pro Thr Gly Gln His	287
ATC CGG TAC CCG AGC AGT CAG CTG Ile Arg Tyr Pro Ser Ser Gln Leu 105	335
AGC CAG TGT GAA TGC AGA CCT AAA Ser Gln Cys Glu Cys Arg Pro Lys 120	383

Fig. 2(ii)

SUBSTITUTE SHEET (RULE 26)

384	AAA Lys	AAG Lys	GAC Asp 130	Ser	GCT Ala	GTG Val	AAG Lys	CCA Pro 135	
432	CGT Arg	CCC Pro 145	CAG Gln	CCC Pro	CGT Arg	TCT Ser	GTT Val 150	CCG Pro	
480	CCC Pro 160	TCC Ser	CCA Pro	GCT Ala	GAC Asp	ATC Ile 165	ACC Thr	CAT His	
528	GCC Ala	CAC His	GCT Ala	GCA Ala	CCC Pro 180	AGC Ser	ACC Thr	ACC Thr	
576	GCT Ala	GCC Ala	GCT Ala	GCC Ala 195	GAC Asp	GCC Ala	GCA Ala	GCT Ala	

Fig. 2(iii)

1									
GAC Asp	AGG Arg	GCT Ala	GCC Ala	ACT Thr 140	Pro	CAC His	C CAC		431
GGC Gly	TGG Trp	GAC Asp	TCT Ser 155	GCC Ala	CCC Pro	GGA Gly	GCA Ala		479
CCC Pro	ACT Thr	CCA Pro 170	GCC Ala	CCA Pro	GGC Gly	CCC Pro	TCT Ser 175		527
Ser	GCC Ala 185	CTG Leu	ACC Thr	CCC Pro	GGA Gly	CCT Pro 190	GCC Ala		575
TCC Ser 200	TCC Ser	GTT Val	GCC Ala	Lys	GGC Gly 205	GGG Gly	GCT Ala	T	624

Fig. 2(iv)

625	AGAGCTCAAC CCAGACACCT GCAGGTGCCG
685	GACTCAGCAG GGTGACTTGC CTCAGAGGCT
745	GGTAAAAAAC AGCCAAGCCC CCAAGACCTC
805	GCCTCTCAGA GGGCTCTTCT GCCATCCCTT
865	GAGTTGGAAG AGGAGACTGG GAGGCAGCAA
825	GGAGTACTGT CTCAGTTTCT AACCACTCTG
985	CTCCCCTCAC TAAGAAGACC CAAACCTCTG
1045	CTGTGACCCC CAACCCTGAT AAAAGAGATG

Fig.2(v)

GAAGCTGCGA AGGTGA	CACA TGGCTTTTC	A 684
ATATCCCAGT GGGGGA	ACAA AGGGGAGCC	r 744
AGCCCAGGCA GAAGCT	GCTC TAGGACCTG	G 804
GTCTCCCTGA GGCCAT	CATC AAACAGGACA	A 864
GAGGGGTCAC ATACCA	GCTC AGGGGAGAA1	924
TGCAAGTAAG CATCTT	ACAA CTGGCTCTTC	984
CATAATGGGA TTTGGGG	CTTT GGTACAAGAA	1044
GAAGGAAAAA AAAAAA	AAA	1094

Fig.2(vi)

14/52	15/52
Fig. 3(i)	Fig.3(ii)

>VEGF_HUMAN VEGF_HUMAN VASCULAR ENDOTHELIAL (VASCULAR 215 AA. LENGTH = 215

SCORE = 181 (92.4 BITS), EXPECT = 6.4e-20, IDENTITIES = 33/75 (44%), POSITIVES = 48/75

QUERY: 31 HQRKVVSWIDVYTRATCQPREVVVPLTVEL

+++ VV +DVY R+ C+P E +V + E

SBJCT: 36 NHHEVVKFMDVYQRSYCHPIETLVDIFQEY

QUERY: 91 PTGQHQVRMQILMIR 105

PT + MQI + I +

SBJCT: 96 PTEESNITMQIMRIK 110

SCORE = 76 (38.8 BITS), EXPECT = 0.0011, IDENTITIES = 12/19 (63%), POSITIVES = 16/19

QUERY: 110 QLGEMSLEEHSQCECRPKK 128

++GEMS +H+ CECRPKK

SBJCT: 116 HIGEMSFLQHNKCECRPKK 134

SCORE = 72 (36.8 BITS), EXPECT = 0.0046, IDENTITIES = 14/21 (66%), POSITIVES = 15/21

QUERY: 202 RCQGRGLELNPDTCRCRKLRR 222

RC +R LELN TCRC K RR

SBJCT: 195 RCKARQLELNERTCRCDKPRR 215

SCORE = 46 (23.5 BITS), EXPECT = 47., IDENTITIES = 6/10 (60%), POSITIVES = 9/10

QUERY: 187 DPRTCRCRCR 196

DP+TC+C C+

SBJCT: 181 DPQTCKCSCK 190

Fig.3(i)

SUBSTITUTE SHEET (RULE 26)

GROWTH FACTOR PRECURSOR (VEGF)

$$P = 6.4e-20$$
 (64%)

MGTVAKQLVPSCVTVQRCGGCCPDDGLECV 90 + PSCV + RCGGCC D+GLECV PDEIEYIFKPSCVPLMRCGGCCNDEGLECV 95

POISSON
$$P(2) = 9.1e-12$$
 (84%)

POISSON
$$P(3) = 3.6e-18$$
 (71%)

POISSON
$$P(4) = 7.3e-10$$
 (90%)

Fig. 3(i)

17/52	18/52
Fig.4(i)	Fig.4(ii)
19/52	20/52
Fig.4(iii)	Fig.4(iv)

Length We Qua R Percen	ight:3.00 ight:0.100 lity:100.9 atio:0.175 t rity:69.703	Average Match:1.000 Average Mismatch:-0.900 Length:739 Gaps:30 Percent Identity:69.703
28	ATGAGCC	CTCTGCTCCGCCGCCTGC
17	 ATGAACTI	TTCTGCTGTCT
68	TGCAGCTC	GCCCCGCCCAGGCCCC
57	TGCTGCTC	TACCTCCACCATGCCAA
118	CACCAGAG	GA
106	 AGAAGGAG	GAGGGCAGAATCATCAC
140	GTGTATAC	TCGC.GCTACCTGCCAG
152	 GTCTATCA	GCGCAGCTA.CTGCCAT
194	TGA.	CTGTGGAGCTCAT
201	TCCAGGAG	
235	CCCAGCTG	CGTGACTGTGCAGCGCT
239	 CCATCCTGT	GTGCCCCTGATGCGAT
285	CCTGGAGTG	TGTGCCCACTGGGCAG
289	CCTGGAGTG	TGTGCCCACTGAGGAG

Fig.4(i)

TGCTCGCCGCACTCC	67
TGGGTGCATTGGAGCCTTGCCT	56
TGTCTCCCAGCCTGATGCCCCTGGC	117
GTGGTCCCAGGCTGCA.CCCATGGC	105
AAGTGGTGTCATGGATAGAT	147
GAAGTGGTGAAGTTCATGGAT	151
CCCCGGGAGGTGGTGGTGCCCT	193
CCAATCGAGACCCTGGTGGACATCT	200
GGGCACCGTGGCCAAACAGCTGGTG	234
GTACATCTTCAA	238
GTGGTGGCTGCCCTGACGATGG	284
GCGGGGCTGCTGCAATGACGAGGG	288
CACCAAGTCCGGATGCAGAT	329
TCCAACATCACCATGCAGATTATGC	338

Fig. 4(ii)
SUBSTITUTE SHEET (RULE 26)

330	CCTCATGATCCGGTACC
339	GGATCAAACCTCAC
369	GTCCCTGGAAGAACACAGCCAGTGT
376	GAGCTTCCTACAGCACAACAATGT
419	GTGCTGTGAAGCCAGACAGGGCTGC
423	GAGCAAGACAAG
469	CGTTCTGTTCCGGGCTGGGACTCTG
443	TGTGGGCCTTGCTCAGA
519	CATCACCCATCCCACTCCAGCCCCA
468	••••••
569	GCACCACCAGCGCCC
469	GCATTTGTTTGTACAA
609	TGCCGACGCCGCAGCTTCCTCCGTT
509	TG.CAAAAACACAGACTCGCGTT
657	AACCCAGACACCTGCAGGTGCCGGA
554	AACGAACGTACTTGCAGATGTGACA
	Fig.4(iii)

	 .
CGAGCAGTCAGCTGGGGGAGAT	368
CAAGGCCAGCACATAGGAGAGAT	375
GAATGCAGACCTAAAAAAAAGGACA	418
GAATGCAGACCAAAGAAAGATA	422
CACTCCCCACCACCGTCCCCAGCCC	468
····· AAAATCCC	442
CCCCCGGAGCACCCTCCCCAGCTGA	518
GCGGAGAA	467
GGCCCCTCTGCCCACGCTGCACCCA	568
· · · · · · ·	468
TGACCCCGGACCTGCCGC	608
GATCCGCAGACGTGTAAATGTTCC	508
GCCAAGGGCGGGCTTAGAGCTC	656
GCAAGGCGAGGCAGCTTGAGTTA	553
AGCTGCGAAGGTGA	695
AGCCGAGGCGGTGA	592

Fig.4(iv)

SUBSTITUTE SHEET (RULE 26)

22 /52	22./22	
22/52	23/52	24/52
Fig.5(i)	Fig.5(ii)	Fig.5(iii)
25/52	26/52	27/52
Fig.5(iv)	Fig.5(v)	Fig.5(vi)

165SOMSQ.MSF.msf MSF:687 Type: D Tuesday, June 20, 1995 Check: 3140 1 VEGF165 ATGAACTTTCTGCTGTCTTGGGTG SOM175 ATGAGCCCTCTGCTCCGCCGCCTG SOM175-e6 ATGAGCCCTCTGCTCCGCCGCCTG SOM175-e6&7 ATGAGCCCTCTGCTCCGCCGCCTG SOM175-e4 ATGAGCCCTCTGCTCCGCCGCCTG 81 VEGF165 CACCCATGGCAGAAGGAGGAGGGC SOM175 TGCCCCTGGCCACCAGAGGAAAGT SOM175-e6 TGCCCCTGGCCACCAGAGGAAAGT SOM175-e6&7 TGCCCCTGGCCACCAGAGGAAAGT SOM175-e4 TGCCCCTGGCCACCAGAGGAAAGT 161 VEGF165 CCAATCGAGACCCTGGTGGACATC SOM175 GTGGTGCCCTTGACTG.TGGA SOM175-e6 GTGGTGCCCTTGACTG.TGGA SOM175-e6&7 GTGGTGCCCTTGACTG.TGGA SOM175-e4 GTGGTGCCCTTGACTG.TGGA 241 VEGF165 GATGCGATGCGGGGGCTGCTAA SOM175 GCAGCGCTGTGGTGGCTGCTCC SOM175-e6 GCAGCGCTGTGGTGGCTGCCC SOM175-e6&7 GCAGCGCTGTGGTGGCTGCCC SOM175-e4 GCAGCGCTGTGGTGGCTGCTCC

Fig.5(i)

CATTGGAGCCTTGCCTTGCTGCTCTACC CTGCTCGCCGCACTCCTGCAGCTGGCCC CTGCTCGCCGCACTCCTGCAGCTGGCCC CTGCTCGCCGCACTCCTGCAGCTGGCCC CTGCTCGCCGCACTCCTGCAGCTGGCCC

AGAATCATCACGAAGTGGTGAAGTTCAT GGTGTCATGGATAGATGTGTATACTCGC GGTGTCATGGATAGATGTGTATACTCGC GGTGTCATGGATAGATGTGTATACTCGC GGTGTCATGGATAGATGTGTATACTCGC

TTCCAGGAGTACCCTGATGAGATCGAGT GCTCATGGGCACCGTGGCCAAAC..AGC GCTCATGGGCACCGTGGCCAAAC..AGC GCTCATGGGCACCGTGGCCAAAC..AGC GCTCATGGGCACCGTGGCCAAAC..AGC

TGACGAGGCCTGGAGTGTGTGCCCACT TGACGATGGCCTGGAGTGTGTGCCCACT TGACGATGGCCTGGAGTGTGTGCCCACT TGACGATGGCCTGGAGTGTGTGCCCACT TGACGATGGCCTGGAGTGTGTGCCCACT

Fig.5(ii)

SUBSTITUTE SHEET (RULE 26)

80)
TCCACCATGCCAAGTGGTCCCAGGCTG	Ì.
CCGCCCAGGCCCTGTCTCCCAGCCTG	βA
CCGCCCAGGCCCTGTCTCCCAGCCTG	λ
CCGCCCAGGCCCTGTCTCCCAGCCTG	ίA
CCGCCCAGGCCCTGTCTCCCAGCCTG	ŀΑ
16	0
GGATGTCTATCAGCGCAGCTACTGCCA	T
GCTACCTGC.CAGCC.CCGGGA	G
24	0
ACATCTTCAAGCCATCCTGTGTGCCCCC	\mathbf{T}
TGGTGCCCAGCTGCGTGACTG	\mathbf{T}
TGGTGCCCAGCTGCGTGACTG	Т
TGGTGCCCAGCTGCGTGACTG	\mathbf{T}
TGGTGCCCAGCTGCGTGACTG	T
32	0
GAGGAGTCCAACATCACCATGCAGATTA	A
GGGCAGCACCAAGTCCGGATGCAGATC	C
GGGCAGCACCAAGTCCGGATGCAGATC	C
GGGCAGCACCAAGTCCGGATGCAGATCC	_
GGGCAGCACCAAGTCCGGATGCAGA	_
	-

Fig.5(iii)

	321
VEGF165	TGCGGATCAAACCTCACCAAGGCC
SOM175	TCATGATCCGGTACCCGAGCA
SOM175-e6	TCATGATCCGGTACCCGAGCA
SOM175-e6&7	TCATGATCCGGTACCCGAGCA
SOM175-e4	••••••••••
	401
VEGF165	AAGAAAGATAGAGCAA
SOM175	AAAAAGGACAGTGCTGTGAAGCCA
SOM175-e6	AAAAAGGACAGTGCTGTGAAGCCA
SOM175-e6&7	AAAAAGGACAGTGCTGTGAAGCCA
SOM175-e4	AAAAAGGACAGTGCTGTGAAGCCA
	481
VEGF165	
SOM175	CTCTGCCCCGGAGCACCCTCCCC
SOM175-e6	• • • • • • • • • • • • • • • • • • • •
SOM175-e6&7	• • • • • • • • • • • • • • • • • • • •
SOM175-e4	CTCTGCCCCGGAGCACCCTCCCC
	561
VEGF165	AGATCCGCA
SOM175	GCACCACCAGCGCCCTGACCCCCG
SOM175-E6	GCACCACCAGCGCCCTGACCCCCG
SOM175-e6&7	GCNCCACCAGCGCC1GACCCCCG
SOM175-e4	GCACCACCAGCGCCCTGACCCCCG
2011273 04	OCACCACCAGCGCCC IGACCCCCG
٠	641
VEGF165	TTGAGTTAAACGAACGTACTTGCA
SOM175	TAGAGCTCAACCCAGACACCTGCA
SOM175-e6	TAGAGCTCAACCCAGACACCTGCA
SOM175-e6&7	•••••••
SOM175-e4	TAGAGCTCAACCCAGACACCTGCA
	Fig.5(iv)
	· · ¬·~ ı··/

STIRSTITITE SHEET (Pula 26)

AGCACATAGGAGAGATGAGCTTCCTACA GTCAGCTGGGGGAGATGTCCCTGGAAGA GTCAGCTGGGGGAGATGTCCCTGGAAGA GTCAGCTGGGGGAGATGTCCCTGGAAGA
GACAAGAAAATCCCTGTGG GACAGGGCTGCCACTCCCCACCACCGTC GATAG
AGCTGACATCACCCATCCCACTCCAGCC
GACGTGTAAATGTTCCTGCAAAAAC.AC GACCTGCCGCTGCCGCTGCCGACGCCGC GACCTGCCGCTGCCGCTGCCGACGCCGC
GATGTGACAAGCCGAGGCGGTGA GGTGCCGGAAGCTGCGAAGGTGA GGTGCCGGAAGCTGCGAAGGTGA .GTGCCGGAAGCTGCGAAGGTGA GGTGCCGGAAGCTGCGAAGGTGA Fig.5(v)

SUBSTITUTE SHEET (Rule 26)

400
GCACAACAAATGTGAATGCAGACCA
ACACAGCCAGTGTGAATGCAGACCTAAA
ACACAGCCAGTGTGAATGCAGACCTAAA
ACACAGCCAGTGTGAATGCAGACCTAAA
······································
TO THE TOTAL CONTRACT OF THE TOTAL CONTRACT
480
·····GCCTTGCTCAGAGCGGAGA
CCCAGCCCCGTTCTGTTCCGGGCTGGGA
CCCAGCCCCGTTCTGTTCCGGGCTGGGA
- COULTETOTICEGGGC IGGGA
560
TTTGTTTGTACA
CCAGGCCCTCTGCCCACGCTGCACCCA
CCAGGCCCTCTGCCCACGCTGCACCCA
CCAGGCCCCTCTGCCCACGCTGCACCCA
The state of the s
640
640 AGACTCGCGTTGCAAGGCGAGGCAGC
AGCTTCCTCCCTTGCAAGGCGAGGCAGC
AGCTTCCTCCGTTGCCAAGGGCGGGCT
AGCTTCCTCCGTTGCCAAGGGCGGGGCT
• • • • • • • • • • • • • • • • • • • •
AGCTTCCTCCGTTGCCAAGGGCCGCCCC

Fig.5(vi)

Fig 6(i)	29/52	F	-ig 6(ii)	30/52
Fig 6(iii)	31/52			

PD ND ·H XX	DA ZA ·H ·J
A A U U A	AA OO · H OH
4 · 00 · 4 0>	4 · 00 · d 0 8
O · D D · F R V	O · Q Q · F F R
 α ⋅ α ∪ α 	α · α α α α α α α α α α α α α α α α α
B · OO AA HA	B · OO AA KKS
KO KK KK KA	KO KK KK DU
A A D D D H A	AA MO DD WK.D
HU UV AU	TCC GC CF
<u> </u>	HA OF ·X XX.
PH < < < HT	
AU DO VA HA	YOU OO A FIELD
H	N N N N N
다 다 다 다 다 다 다 다 다 다 다	
14 X> ·X 4D	DD X CX PL
44 FT XX XD	AA FI XX OD
HO KK DU	HO KK D ·
S · Y X D D R F	α · > × ф ф ф .
H A A A A A A A A A A A A A A A A A A A	L RR PE·S
HU NO UK	H H O O H K
マ コ 田 田 田 田 田 田 田	> h B H B H A Q
BK DO DO NE	BK CO CO KH
SK PM KO KE	SH PM MAH
NO NZ LK	DH SZ LK LL
THE EN EN EN EN EN EN EN EN EN EN EN EN EN	TH EE HH OF
FO GO GE	FO QD QE
No Fe TE XA	NO FE TH PK
HO LH KK	DE LT FT DG
rt rt	
Sho	ong page
165 75 75 75 75 75	$\frac{65}{5}$
GF 3F 11 7 11 7 11 7 11 7 11 7 11 7 11 7	F1 17 17 17 17 17 17 17 17 17 17 17 17 17
VEGF ₁₆₅ SOM175 _{Short} VEGF ₁₆₅ SOM175 _{Short} VEGF ₁₆₅ SOM175 _{Short} VEGF ₁₆₅ SOM175 _{Short}	VEGF ₁₆₅ SOM175 _{Long} VEGF ₁₆₅ SOM175 _{Long} VEGF ₁₆₅ SOM175 _{Long} VEGF ₁₆₅ SOM175 _{Long}
	VEGF ₁₆₅ SOM175 _{Long} VEGF ₁₆₅ SOM175 _{Long} VEGF ₁₆₅ SOM175 _{Long} VEGF ₁₆₅ SOM175 _{Long}
SIRSTITE	

			5	30/5	2				
60	121 115	170 175	191 207		60 55		1 / 1	, 62	
Дд	က က	ΕÆ			ДД	4 00 0			
>>		S S	_		>>			_	
니>		다 다			I >	Б	i . p	4	
E >	_	D O			E >	<u> </u>	ی ۰ را) :	
田田田	,	Ο̈́			田田] H 🗗	1 · D	,	rlg.o(11,
HR	OH OH OH	> 4			HR	_			J. C
C P	J	• Д,			<u></u> 라	J		į	<u> </u>
[D D]	D C D C	L F T			HO	_	_		
y F	, b H K	HH			00	•			
SA	Д,	X F			S Y A T	H d	· II		
民民	X X	жн			医民民		· [+		
Q F	НН	R D		•	Q FI	HH	н · О · [民民	
KK	K Z	田及			K K	K Z	. A	N N H	
>>	Z l	വ പ			>>	E K	· A		
	нн	ညက				НН	• 03	X X	
ΣH	0 0	다 다			ΣH	00	• Д	DA	
F N	ΣΣ	U A			Fi 🔀	ZZ	· 4	CC	
X Q	FIK	ည			X W	EHK	. ტ	2 2	
	H >	요 라			>>	H >	• д	00	
<u>> ></u>	ZO	ZK			> >	ZQ	· 4		
田民	S H	田の			· X	SH	· W	K D	
HHO	田 G 田	Q D			田民	E O	· D	되 다	
HH	E E	. Q ⊠ №			HO	E O	· 3	ZZ	
N U	<u> </u>	• д			HH	E E	· છ	나 다	
O Di	> >	• >			Z O	<u>Ч</u>	. ሷ	田田	
U A	00	• W			A P P	> >	. >	디디	
O D	回田	· ¤			о С	E E	. Ω • Ω	Q D	
ር ር	디디	• д			о С	그기	N P R	K K	
E E	<u>ი</u> ი	· 0	M A	•	ыO	0 0	E O	X Q G A	
A W	HО	• д	よ ら		A W	田口	D G		
Z >	ΩΩ	· K	ው ው		Σ>	ДΩ	民民	R R <t< td=""><td></td></t<>	
							i		

SURSTITUTE OUTER IN THE

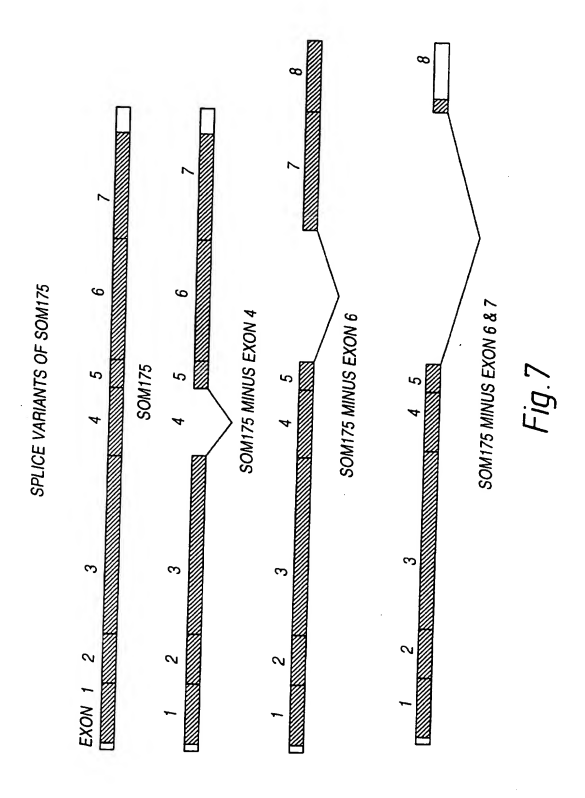
Areas of 100% homology are boxed and conserved residues thought sequence (removal of which gives rise to mature VEGF₁₆₅) giving The VEGF sequence depicted includes the 26 amino acid leader to be involved in homodimerisation are underlined. total length of 191 amino acids.

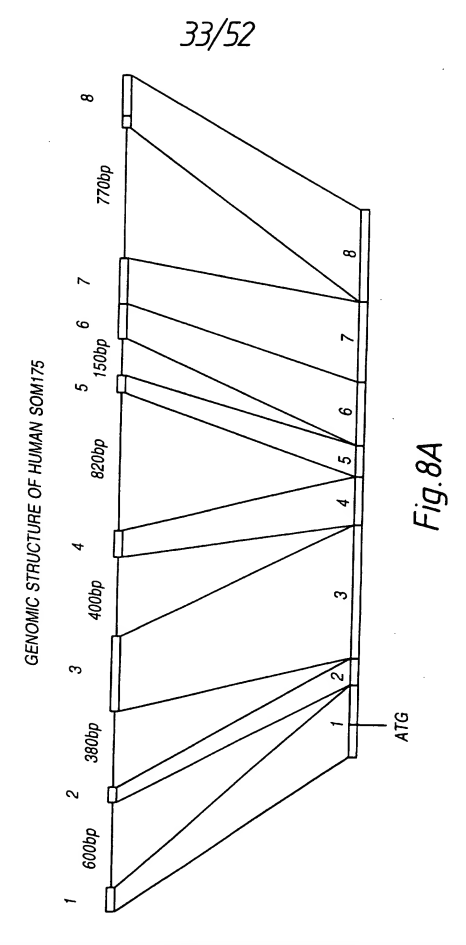
82 including those thought to be involved in homodimerisation Arginine-77, Cystein-78, Glycine-80, Cysteines-81 Homology of SOM175 to $VEGF_{165}$ is 27% (33%) at the protein level, however within this are blocks of 100% homology. In particular, many structural residues are conserved Proline-70, Cysteine-72, Valine-74 of VEGF (by comparison with PDGF). Cysteine-47 SUBSTITUTE SHEET (RULE 26)

Fig.6 (iii)

Cysteine-89, Proline-91

Cysteines 122 & 124





SUBSTITUTE SHEET (Rule 26)

			34	-/52	•		
GGCCAG gtacgtgagg	aatacttaca	ATGCAG gtccgagatg	ATGCAG gtgtcaggca	AGACAG gtgagtcttt	cccaggccc	acacctgtag	3'UTR
GGCCAG	GGAAAG	ATGCAG	ATGCAG	AGACAG	CTCCAG	ACCCAG	AGG TGA
(dq09)	(43bp)	(187bp)	(73bp)	(34bb)	Exon 6 (101bp)	(109bp)	(22bp)
*Exon 1	Exon 2	Exon 3	Exon 4	Exon 5	Exon 6	Exon 7	*Exon 8
ATGAGG	GCCCCT	TGGTGT	ATCCTC	ACCTAA	GGCTGC	CCCCAG	GTGCCG
5'UTR	tctcccacag	tctgctccca	ctgaatacag	acttttcaag	ctcctccgta GGCTGC	cccactccag	ccctgctcag GTGCCG

Fig.8B

36/52	37/52
Fig. 9(i)	Fig. 9(ii)
. 38/52	3 <u>9</u> /52
Fig.9(iii)	Fig. 9(iv)

-163 -103 -43	g g	ggg	rccg	rcgg	ragg	ago	cgc	ccc	cta	gcto cgco acco
16									GCT	GCAG
	R	R	L	L	L	V	A	L	L	Q
76	${f T}{f T}'$	TGA'	TGG	CCC	CAG!	rca(CCA	GAA	GAAZ	↓ AGTG
	F	D	G	Р			Q	K	K	V
136	AC	ልጥር (ירא(3000	~ > ~ ~ ~	יר אר	~~m/	2000		CCT
	T	C	O	P		GA(E	V V	3G'I'() V	GTG V	
			Z.	•	10		V	V	V	P
196			CTA	AGT(GCCC	'AGC	TGI	GTG	ACT	GTG
	K	Q	L	V	P	S	С	V	${f T}$	V
256	GGC	'CTG	GAA	ጥርባ	ነርጥር	CCC	י א רית	יררה	CAA	
	G	L	E	C		P	T	G	O	H
216									~	
316	TAC						GGG	GAG.	ATG:	rcc
	Y	P	S	S	Q	L	G	E	M	S
376	CCT.	AAA	AAA	AAG	GAG	AGT(GCT(GTG:	ACC	ר א
	P	K	K	K	E	S	A	V	R	P
136	C 3 C 4	700	200-							
±30	CAGO								ACT	CT
	Q	P	R	S	V	P	G	W	D	S

Fig.9(i)

Ī				_	_			_	ggga	
								-	atgg	
CC	ggg	ctag	ggg	ccc	TAg	<u>a</u> AG(CCC	CCTC	GCTG	
					M	S	P	L	L	-17
					1					
CTC	GGC	rcgo	CAC	CCAC	GCC	CCC	rgro	STCC	CAG	
L	Α	R	${f T}$	0	A		V	S	0	4
				~					~	_
GTO	GCCA	ATGO	ATA	AGAC	CGTT	TAT	rgca	CGT	GCC	
V	P	W	I	D	V	Y	A	R	A	24
			_		•	_			**	2 -
CTC	AGC	ATG	GAA	CTC	'ATC	GGC	ΊΑΑΤ	ነርጥር	GTC	
L	S	M	E	L	M	G	N	V	V	44
			_		••	J	-1	٧	V	* *
CAG	ירהר	יጥርጥ	יככיד	יככר	יחיבר	ጥርር	ירכתי	GAC	GAT	
0	R	C			C			D	D	6.1
Ž	11	C	G	G	1	C	P	ע	D	64
$C \lambda \lambda$	СШС		א חור		A MC	ama	7 m	3 M.C	~ ~ ~	
							ATG			
Q	V	R	M	Q	I	L	M	I	Q	84
ama		~~~	~~	. ~ ~	~ ~ ~					
							GAA			
L	G	E	H	S	Q	C	E	C	R	104
	•									
GAC	AG <u>G</u>	<u>GTT</u>	<u>GCC.</u>	ATA	CCC	CAC	CAC	CGT	CCC	
. D	R	V	Α	I	P	H	Η	R	P	124
<u>ACC</u>	CCG	GGA(GCA	CCC'	TCC(CCA	GCT(<u>GAC</u>	ATC	
${f T}$	P	G	A	P	S	P	A	D	I	144

Fig.9(ii)

496	7 m.c	1 (7 N III)	~~~	3 O.F			~~~			
430	ATC	.CAT	CCC	AC'I	'CCZ	<u>AG</u> C	CCC	AGG	ATC	CTCT
	I	H	P	${f T}$	P	A	P	G	S	S
						S	P	R	I	L
556	CTG	ACC	CCC	GGA	.CCI	rgco	CGT	rgc	CGC!	ГGТА
	L	${f T}$	Р	G	P	A	V	Α	A	V
	P	D :	P]	R	\mathbf{T}	С	R	С	R	С
C1 C	~~~	~~								
616				AGC	TCA	ACC	CAC	SACA	ACC.	rgta
	G	A	*							
	R	G I	٠	Ξ :	L	N	P	D	T	С
676	ctt	tcca	agag	rtc	cac	יממט	, C C C	יממכ	·+~	ttt.
736									_	
										ıgag
796	gag	ctct	cctc	gc	cat	ctt	tta	tct	CCC	aga
856	atg	tctc	cacc	ctca	agg	ggc	cag	ggt	act	ctc
916	ttc						_			
976	gggt									
1036	gaca							•		
	9400	~ C C C	·uuu	uuc	ıua	uaa	uaa	uaa	aaa	aaa

Fig.9(iii)

								—	-			
	GC	CCG	CCጥ	ጥርር	ልሮሮ	C A C	CCC	CCC	ሮ አ አ	CGCC		
	A	R		A	ACC P					_		
	C	P	P		_	S	_A	_A	N	A	164	;
		P	P	С	\mathbf{T}	Q	R	R	Q	R	130	1
	C Λ (~~~		000	TIES C.	ama.	~					
	_		_			CTC	CA'I"	I'GC(CAA	GGGC		
	_ D	A	A	A	S	S	I	A	K	G	184	
	R	R	R	R	F	L	H	C	Q	G	150)
	1											
	GG1	rgc	CGGZ	AAG	CCG	CGAA	\AG <u>'</u>	<u>rga</u>	caag	gctg		
											186	
	R	C	R	K	P	R	к *	r			167	
i											1. 0 7	
	tat	ggd	ccct	gct	tca	cao	aaa	gaa	gac	ıtgg		
	gto	act	acc	cca	laas	act	aas	cct	+++	aga		
	act	acc	ato	taa	CAA	tta	tca	one.	220	ctc		
	tca	ctt	aac	cac	cct	aat	Caa	.agg	200	atc		
	aac			att	a+ 5	ggc	caa	.g cg	ayo	atc		
_	200	===	laaa IL		.CLa	.cca	ala	acg	gga	ttt		٠
	aca					gat	aaa	aga	gat	gga		
	aaa	aaa	aaa	aaa								

Fig.9(iv)

41/52	42/52
Fig 10(i)	Fig 10(ii)

A		
hVRF167	-21 MSPLLRRLLLAALLQLAPAQA	P
mVRF167		P
hVRF167	30 EVVVPLTVELMGTVAKQLVPS	
mVRF167	: 30 EVVVPLSMELMGNVVKQLVPS(
hVRF167	80 ILMIRYPSSQLGEMSLEEHSQC	
mVRF167		
hVRF167	130 RPDPRTCRCRCRRRSFLRCQGR	2
mVRF167) L
В		
hVRF186	116 RAATPHHRPQPRSVPGWDSAPG	
mVRF186	116 RVAIPHHRPQPRSVPGWDSTPG	
hVRF186	166 TPGPAAAAADAAASSVAKGGA*	
mVRF186	166 TPGPAVAAVDAAASSIAKGGA*	

Fig.10(i)

VSQPDAPGHQRKVVSWIDVYTRATCQPR 	29
VSQFDGPSHQKKVVPWIDVYARATCQPR	29
VTVQRCGGCCPDDGLECVPTGQHQVRMQ	79
VTVQRCGGCCPDDGLECVPTGQHQVRMQ	79
ECRPKKKDSAVKPDSPRPLCPRCTQHHQ	129
ECRPKKKESAVRPDSPRILCPPCTQRRQ	129
GLELNPDTCRCRKLRR* 167	
GLELNPDTCRCRKPRK* 167	
APSPADITHPTPAPGPSAHAAPSTTSAL 1	.65
APSPADIIHPTPAPGSSARLAPSAANAL 1	.65
186	
186	

Fig.10(ii)

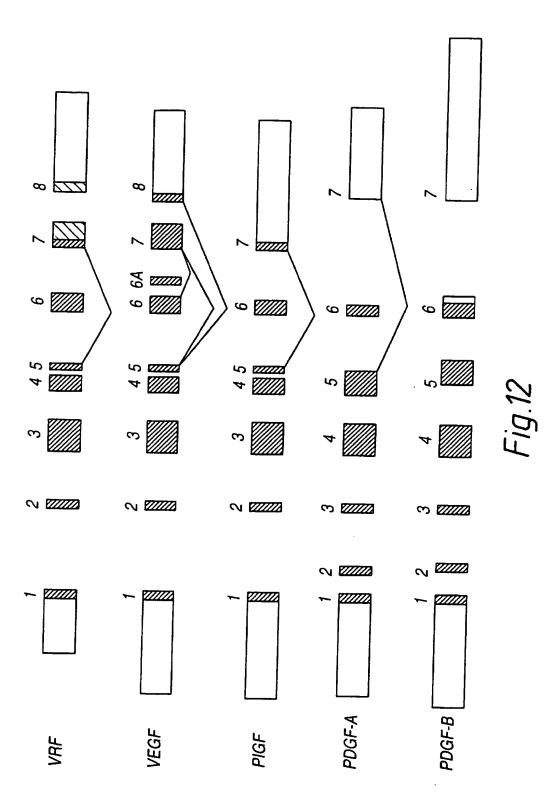
44/52	45/52
Fig 11(i)	Fig 11(ii)

mVRF167	-21 MSPLLRRLLLVALLQL
mVEGF188	:: : -26 MNFLLSWVHWTLALLLYLHH
mVRF167	25 TCQPREVVVPLSMELMGNVV
mVEGF188	
mVRF167	75 QVRMQILMIQYPSSQ.LGEM
mVEGF188	: : : 74 NITMQIMRIKPHQSQHIGEM
mVRF167	119ILCPPC
mVEGF188	: 124 QKRKRKKSRFKSWSVHCEPC
mVRF167	152 GLELNPDTCRCRKPRK
mVEGF188	: 173 QLELNERTCRCDKPRR

Fig.11(i)

1	
AR.TQAPVSQFDGPSHQKKVVPWIDVYARA	24
AKWSQAAPTT.EGEQKSHEVIKFMDVYQRS	23
KQLVPSCVTVQRCGGCCPDDGLECVPTGQH : : : : : :	74
YIFKPSCVPLMRCAGCCNDEALECVPTSES	73
SLGEHSQCECRPKKKESAVRPDSPR	118
SFLQHSRCECRPKKDRTKPEKKSVRGKGKG	123
TQRRQRPDPRTCRCRCRRRRFLHCQGR : : : : :	151
SERRKHLFVQDPQTCKCSCKNTDS.RCKAR	172
	167
	188

Fig.11(ii)



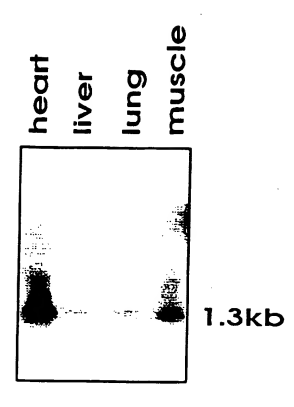


Fig.13

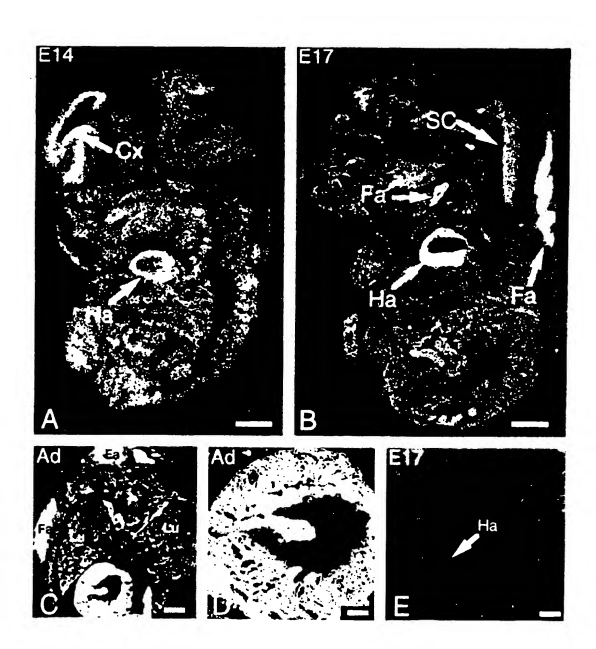
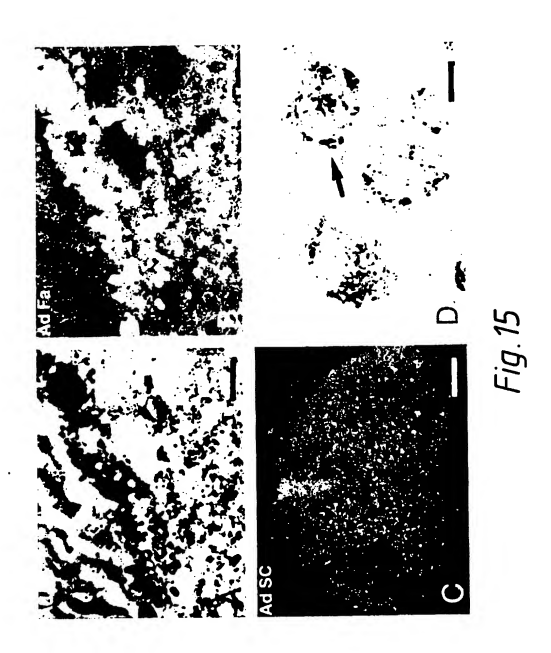
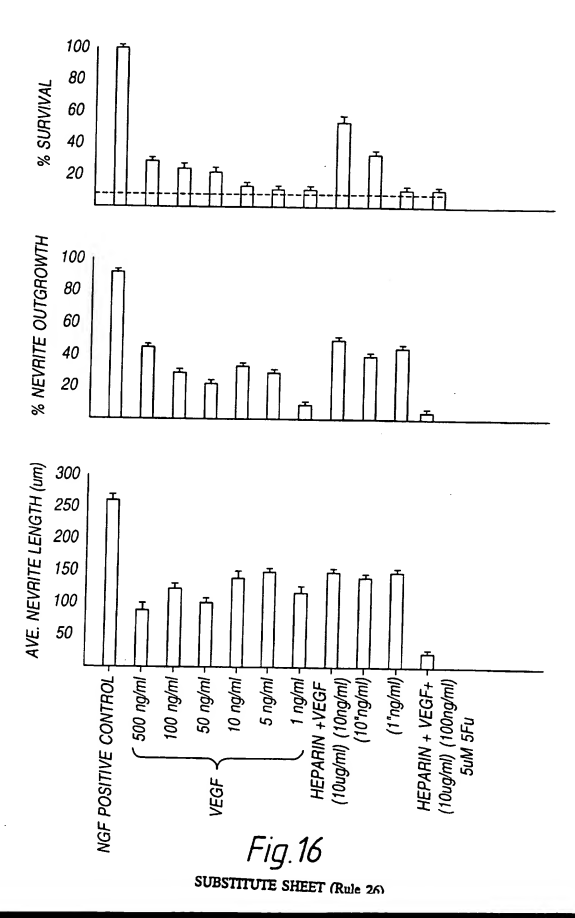
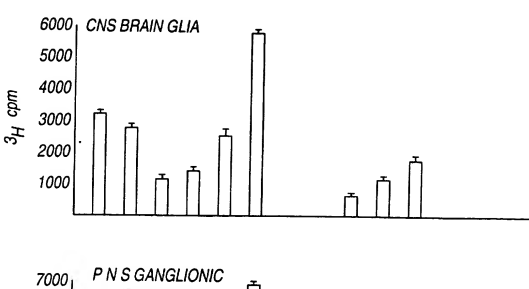
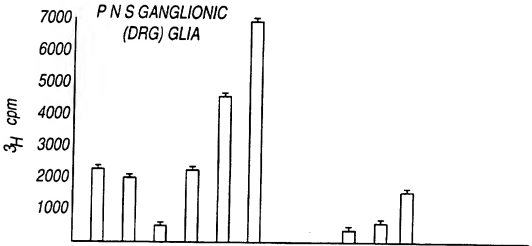


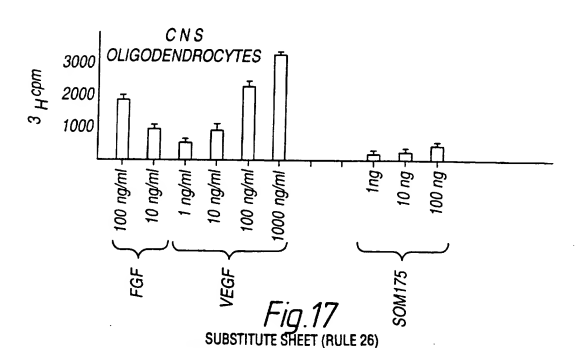
Fig.14







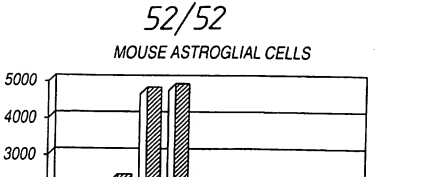




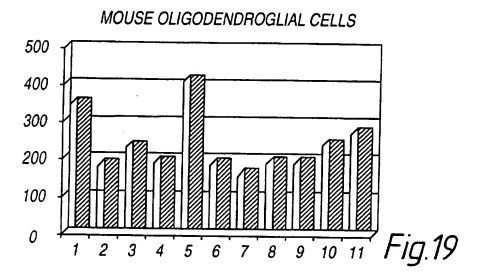
2000

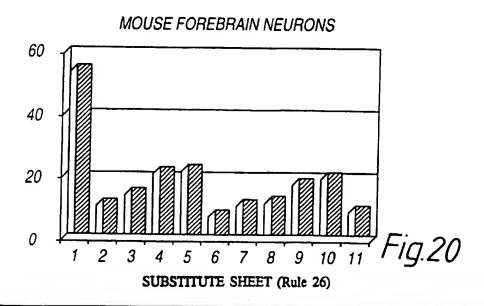
1000

0



2 3 4 5 6 7 8 9 10 11 Fig.18





INTERNATIONAL SEARCH REPORT

International Application No. PCT/AU 96/00094

A. CLASSIFICATION OF SUBJECT MATTER

Int Cl6: C12N 15/12; C07K 14/475; A61K 037/02

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) WPAT AND CHEM ABS

SEE DETAILS IN ELECTRONIC DATABASE BOX BELOW

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched USPM, JAPIO

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
DERWENT WPAT, USPM, JAPIO DATABASES; <u>KEYWORDS</u>: MVRF OR HVRF OR SOM 1: OR SOM X:
OR [VASOACTIVE () PERMEABILITY () FACTOR#] OR VEGF: OR VEGF OR VRF: OR VRF OR
[GROWTH () FACTOR (5N) (VASCULAR OR ENDOTHELI:]
CHEMICAL ABSTRACTS DATABASE; KEYWORDS: AS ABOVE

C. DOCUMENTS CONSIDERED TO BE RELEVANT Category* Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. X AU 60798/90 (CALIFORNIA BIOTECHNOLOGY INC) published 1-41 21 February 1991 X AU 56574/90 (GENENTECH, INC.) published 15 November 1990 1-41 P,X AU 73941/94 (HUMAN GENOME SCIENCES, INC.) published 1-41 14 September 1995 Further documents are listed in the continuation of Box C X See patent family annex Special categories of cited documents: "T" later document published after the international filing date or "A" document defining the general state of the art which is priority date and not in conflict with the application but cited to not considered to be of particular relevance understand the principle or theory underlying the invention "E" earlier document but published on or after the document of particular relevance; the claimed invention cannot international filing date be considered novel or cannot be considered to involve an "L" document which may throw doubts on priority claim(s) inventive step when the document is taken alone or which is cited to establish the publication date of document of particular relevance; the claimed invention cannot another citation or other special reason (as specified) be considered to involve an inventive step when the document is "O" document referring to an oral disclosure, use, combined with one or more other such documents, such exhibition or other means combination being obvious to a person skilled in the art •P• document published prior to the international filing document member of the same patent family date but later than the priority date claimed Date of the actual completion of the international search Date of mailing of the international search report 1996. 03 June 1996 5тн JUNE Name and mailing address of the ISA/AU Authorized officer AUSTRALIAN INDUSTRIAL PROPERTY ORGANISATION **PO BOX 200** WODEN ACT 2606 ARATI SARDANA AUSTRALIA Facsimile No.: (06) 285 3929 Telephone No.: (06) 283 2627

INTERNATIONAL SEARCH REPORT

International Application No.

C (Continua	ntion) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to
х	US 5073492 (Chung-Ho Chen and Sumi C. Chen) published 17 December 1991	claim No.
х	Biochemical and Biophysical Research Communications (1992), Vol 183, No. 3, pg 1167-1174 (Weindel K. et al.) "AIDS-ASSOCIATED KAPOSI'S SARCOMA CELLS IN CULTURE EXPRESS VASCULAR ENDOTHELIAL GROWTH FACTOR". See whole Article.	1-41
x	Biochemical and Biophysical Research Communications (1989), Vol 165, No. 3 pg 1198-1206 (Edmund Tischer et al.) "VASCULAR ENDOTHELIAL GROWTH FACTOR: A NEW MEMBER OF THE PLATELET-DERIVED GROWTH FACTOR GENE FAMILY". See whole Article.	1-41
х	Journal of Virology (1994), Vol 68, No. 1 pg 84-92 (David J. Lyttle et al.) "HOMOLOGS OF VASCULAR ENDOTHELIAL GROWTH FACTOR ARE ENCODED BY THE POX VIRUS ORF VIRUS". See whole Article.	1-41
х	Methods in Enzymology (1991), vol 198, pg 391-405 (Ferrara Napoliana et al.) "PURIFICATION AND CLONING OF VASCULAR ENDOTHELIAL GROWTH FACTOR SECRETED BY PITUITARY FOLLICULOSTELLATE CELLS". See whole Article	1-41
х	The Journal of Biological Chemistry (1991), vol 266, No. 18 pg 11947-11954 (Edmund Tischer et al.) "THE HUMAN GENE FOR VASCULAR ENDOTHELIAL GROWTH FACTOR". See whole Article	1-41
P,X	Biochemical and Biophysical Research Communications (1996), Vol 220, No. 1 pg 147-52 (Lagercrantz J et al) "EXPRESSION OF THE VEGF-RELATED FACTOR GENE IN PRE-AND POSTNATAL MOUSE". See whole Article	33
	Biochimica et Biophysica Acta (1995), Vol. 1260 No. 2 pg 235-9 (Sharma Hari S et al.) "NUCLEOTIDE SEQUENCE AND EXPRESSION OF THE PORCINE VASCULAR ENDOTHELIAL GROWTH FACTOR". See whole Article.	1-41
	DEVELOPMENT (1992), Vol 114, pg 521-532 (Breier G et al.) "EXPRESSION OF VASCULAR ENDOTHELIAL GROWTH FACTOR DURING EMBRYONIC ANGIOGENESIS AND ENDOTHELIAL CELL DIFFERENTIATION". See whole Article	1-41
1 5	Molecular Endocrinology (1991), Vol 5 No. 12, pg 1806-1814 (Houck Keith A et al.) "THE VASCULAR ENDOTHELIAL GROWTH FACTOR FAMILY: IDENTIFICATION OF A FOURTH MOLECULAR SPECIES AND CHARACTERIZATION OF ALTERNATIVE SPLICING OF RNA". See whole Article	1-41

INTERNATIONAL SEARCH REPORT

This Annex lists the known "A" publication level patent family members relating to the patent documents cited in the above-mentioned international search report. The Australian Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Patent Do	cument Cited in Search Report	ı		Patent	Family Member		
AU 56574/90	US IL WO	5332671 94128 9013649	CA JP	2054699 4505705	EP NZ	471754 233451	
AU	60798/90	US JP	5194596 5501350	CA WO	2063810 9102058	EP US	484401 5219739
ΑU	73941/94	wo	9524473				
US	5073492	NONE					

END OF ANNEX